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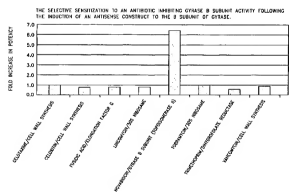
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[Continued on next page]

(54) Title: IDENTIFICATION OF ESSENTIAL GENES IN PROKARYOTES



(57) Abstract: The sequences of antisense nucleic acids which inhibit the proliferation of prokaryotes are disclosed. Cell-based assays which employ the antisense nucleic acids to identify and develop antibiotics are also disclosed. The antisense nucleic acids can also be used to identify proteins required for proliferation, express these proteins or portions thereof, obtain antibodies capable of specifically binding to the expressed proteins, and to use those expressed proteins as a screen to isolate candidate molecules for rational drug discovery programs. The nucleic acids can also be used to screen for homologous nucleic acids that are required for proliferation in cells other than Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, and Pseudomonas aeruginosa. The nucleic acids of the present invention can also be used in various assay systems to screen for proliferation required genes in other organisms.

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IDENTIFICATION OF ESSENTIAL GENES IN PROKARYOTES

Sequence Listing

The present application is being filed along with duplicate copies of a CD-ROM marked "Copy 1" and "Copy 2" containing a Sequence Listing in electronic format. The duplicate copies of the CD-ROM each contain a file entitled SEQLIST_FINAL_9PM created on March 20, 2001 which is 37.487,912 bytes in size.

Background of the Invention

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Since the discovery of penicillin, the use of antibiotics to treat the ravages of bacterial infections has saved millions of lives. With the advent of these "miracle drugs," for a time it was popularly believed that humanity might, once and for all, be saved from the scourge of bacterial infections. In fact, during the 1980s and early 1990s, many large pharmaceutical companies cut back or eliminated antibiotics research and development. They believed that infectious disease caused by bacteria finally had been conquered and that markets for new drugs were limited. Unfortunately, this belief was overly optimistic.

The tide is beginning to turn in favor of the bacteria as reports of drug resistant bacteria become more frequent. The United States Centers for Disease Control announced that one of the most powerful known antibiotics, vancomycin, was unable to treat an infection of the common Staphylococcus aureus (staph). This organism is commonly found in our environment and is responsible for many nosocomial infections. The import of this announcement becomes clear when one considers that vancomycin was used for years to treat infections caused by Staphylococcus species as well as other stubborn strains of bacteria. In short, bacteria are becoming resistant to our most powerful antibiotics. If this trend continues, it is conceivable that we will return to a time when what are presently considered minor bacterial infections are fatal diseases.

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Over-prescription and improper prescription habits by some physicians have caused an indiscriminate increase in the availability of antibiotics to the public. The patients are also partly responsible, since they will often improperly use the drug, thereby generating yet another population of bacteria that is resistant, in whole or in part, to traditional antibiotics.

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The bacterial pathogens that have haunted humanity remain, in spite of the development of modern scientific practices to deal with the diseases that they cause. Drug resistant bacteria are now an increasing threat to the health of humanity. A new generation of antibiotics is needed to once again deal with the pending health threat that bacteria present.

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Discovery of New Antibiotics

As more and more bacterial strains become resistant to the panel of available antibiotics, new antibiotics are required to treat infections. In the past, practitioners of pharmacology would have to rely upon traditional methods of drug discovery to generate novel, safe and efficacious compounds for the treatment of disease. Traditional drug discovery methods involve blindly testing potential drug candidate-molecules, often selected at random, in the hope that one might prove to be an effective treatment for some disease. The process is painstaking and laborious, with no guarantee of success. Today, the average cost to discover and develop a new drug exceeds US \$500 million, and the average time from laboratory to patient is 15 years. Improving this process, even incrementally, would represent a huge advance in the generation of novel antimicrobial agents.

Newly emerging practices in drug discovery utilize a number of biochemical techniques to provide for directed approaches to creating new drugs, rather than discovering them at random. For example, gene sequences and proteins encoded thereby that are required for the proliferation of a cell or microorganism make excellent targets since exposure of bacteria to compounds active against these targets would result in the inactivation of the cell or microorganism. Once a target is identified, biochemical analysis of that target can be used to discover or to design molecules that interact with and alter the functions of the target. Use of physical and computational techniques to analyze structural and biochemical properties of targets in order to derive compounds that interact with such targets is called rational drug design and offers great potential. Thus, emerging drug discovery practices use molecular modeling techniques, combinatorial chemistry approaches, and other means to produce and screen and/or design large numbers of candidate compounds.

Nevertheless, while this approach to drug discovery is clearly the way of the future, problems remain. For example, the initial step of identifying molecular targets for investigation can be an extremely time consuming task. It may also be difficult to design molecules that interact with the target by using computer modeling techniques. Furthermore, in cases where the function of the target is not known or is poorly understood, it may be difficult to design assays to detect molecules that interact with and alter the functions of the target. To improve the rate of novel drug discovery and development, methods of identifying important molecular targets in pathogenic cells or microorganisms and methods for identifying molecules that interact with and alter the functions of such molecular targets are urgently required.

Staphylococcus aureus is a Gram positive microorganism which is the causative agent of many infectious diseases. Local infection by Staphylococcus aureus can cause abscesses on skin and cellulitis in subcutaneous tissues and can lead to toxin-related diseases such as toxic shock and scalded skin syndromes. Staphylococcus aureus can cause serious systemic infections such as osteomyelitis, endocarditis, pneumonia, and septicemia. Staphylococcus aureus is also a common cause of food poisoning, often arising from contact between prepared food and infected food industry workers. Antibiotic resistant strains of Staphylococcus aureus have recently been

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identified, including those that are now resistant to all available antibiotics, thereby severely limiting the options of care available to physicians.

Pseudomonas aeruginosa is an important Gram-negative opportunistic pathogen. It is the most common Gram-negative found in nosocomial infections. P. aeruginosa is responsible for 16% of nosocomial pneumonia cases, 12% of hospital-acquired urinary tract infections, 8% of surgical wound infections, and 10% of bloodstream infections. Immunocompromised patients, such as neutropenic cancer and bone marrow transplant patients, are particular susceptible to opportunistic infections. In this group of patients, P. aeruginosa is responsible for pneumonia and septicemia with attributable deaths reaching 30%. P. aeruginosa is also one of the most common and lethal pathogens responsible for ventilator-associated pneumonia in intubated patients, with directly attributable death rates reaching 38%. Although P. aeruginosa outbreaks in burn patients are rare, it is associated with 60% death rates. In the AIDS population, P. aeruginosa is associated with 50% of deaths. Cystic fibrosis patients are characteristically susceptible to chronic infection by P. aeruginosa, which is responsible for high rates of illness and death. Current antibiotics work poorly for CF infections (Van Delden & Igelwski. 1998. Emerging Infectious Diseases 4:551-560; references therein).

The gram-negative enteric bacterial genus, Salmonella, encompasses at least 2 species. One of these, S. enterica, is divided into multiple subspecies and thousands of serotypes or serovars (Brenner, et al. 2000 J. Clin. Microbiol. 38:2465-2467). The S. enterica human pathogens include serovars Typhi, Paratyphi, Typhimurium, Cholerasuis, and many others deemed so closely related that they are variants of a widespread species. Worldwide, disease in humans caused by Salmonella is a very serious problem. In many developing countries, S. enterica ser. Typhi still causes oftenfatal typhoid fever. This problem has been reduced or eliminated in wealthy industrial states. However, enteritis induced by Salmonella is widespread and is the second most common disease caused by contaminated food in the United States (Edwards, BH 1999 "Salmonella and Shigella species" Clin. Lab Med. 19(3):469-487). Though usually self-limiting in healthy individuals, others such as children, seniors, and those with compromising illnesses can be at much greater risk of serious illness and death.

Some S. enterica serovars (e.g. Typhimurium) cause a localized infection in the gastrointestinal tract. Other serovars (i.e. Typhi and Paratyphi) cause a much more serious systemic infection. In animal models, these roles can be reversed which has allowed the use of the relatively safe S. enterica ser. Typhimurium as a surrogate in mice for the typhoid fever agent, S. enterica ser. Typhi. In mice, S. enterica ser Typhimurium causes a systemic infection similar in outcome to typhoid fever. Years of study of the Salmonella have led to the identification of many determinants of virulence in animals and humans. Salmonella is interesting in its ability to localize to and invade the intestinal epithelium, induce morphologic changes in target cells via injection of certain cell-remodeling proteins, and to reside intracellularly in membrane-bound vesicles (Wallis, TS and

Galyov, EE 2000 "Molecular basis of *Salmonella*-induced enteritis." Molec. Microb. 36:997-1005; Falkow, S "The evolution of pathogenicity in Escherichia, Shigella, and Salmonella," Chap. 149 in Neidhardt, et al. eds pp 2723-2729; Gulig, PA "Pathogenesis of Systemic Disease," Chap. 152 in Neidhardt, et al. ppp 2774-2787). The immediate infection often results in a severe watery diarrhea but *Salmonella* also can establish and maintain a subclinical carrier state in some individuals. Spread is via food contaminated with sewage.

The gene products implicated in Salmonella pathogenesis include type three secretion systems (TTSS), proteins affecting cytoplasmic structure of the target cells, many proteins carrying out functions necessary for survival and proliferation of Salmonella in the host, as well as "traditional" factors such as endotoxin and secreted exotoxins. Additionally, there must be factors mediating species-specific illnesses. Despite this most of the genomes of S. enterica ser. Typhi (see http://www.sanger.ac.uk/Projects/S_typhi/ for the genome database) and S. enterica ser. Typhimurium (see http://genome.wustl.edu/gsc/bacterial/salmonella.shtml for the genome database) are highly conserved and are mutually useful for gene identification in multiple serovars. The Salmonella are a complex group of enteric bacteria causing disease similar to but distinct from other gram-negative enterics such as E. coli and have been a focus of biomedical research for the last century.

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Enterococcus faecalis, a Gram-positive bacterium, is by far the most common member of the enterococci to cause infections in humans. Enterococcus faecium generally accounts for less than 20% of clinical isolates. Enterococci infections are mostly hospital-acquired though they are also associated with some community-acquired infections. Of nosocomial infections enterococci account for 12% of bacteremia, 15% of surgical wound infections, 14% of urinary tract infections, and 5 to 15% of endocarditis cases (Huycke, M. M., D. F., Sahm and M. S. Gilmore. 1998. Emerging Infectious Diseases 4:239-249). Additionally enterococci are frequently associated with intraabdominal and pelvic infections. Enterococci infections are often hard to treat because they are resistant to a vast array of antimicrobial drugs, including aminoglycosides, penicillin, ampicillin and vancomycin. The development of multiple-drug resistant (MDR) enterococci has made this bacteria a major concern for treating nosocomial infections.

These reasons underscore the urgency of developing new antibiotics that are effective against Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, and Enterococcus faecalis. Accordingly, there is an urgent need for more novel methods to identify and characterize bacterial genomic sequences that encode gene products involved in proliferation, and are thereby potential new targets for antibiotic development. Prior to the present invention, the discovery of Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, and Pseudomonas aeruginosa and Enterococcus faecalis genes required for proliferation of the microorganism was a painstaking and slow process. While the detection of new cellular drug targets within a Staphylococcus aureus, Salmonella typhimurium, Klebsiella

pneumoniae, Pseudomonas aeruginosa or Enterococcus faecalis cell is key for novel antibiotic development, the current methods of drug target discovery available prior to this invention have required painstaking processes requiring years of effort.

Summary of the Invention

Some aspects of the present invention are described in the numbered paragraphs below.

1. A purified or isolated nucleic acid sequence comprising a nucleotide sequence consisting essentially of one of SEQ ID NOs: 8-3795, wherein expression of said nucleic acid inhibits proliferation of a cell.

- 2. The nucleic acid sequence of Paragraph 1, wherein said nucleotide sequence is complementary to at least a portion of a coding sequence of a gene whose expression is required for proliferation of a cell.
- 3. The nucleic acid of Paragraph 1, wherein said nucleic acid sequence is complementary to at least a portion of a nucleotide sequence of an RNA required for proliferation of a cell.
- 4. The nucleic acid of Paragraph 3, wherein said RNA is an RNA comprising a sequence of nucleotides encoding more than one gene product.
 - 5. A purified or isolated nucleic acid comprising a fragment of one of SEQ ID NOs.: 8-3795, said fragment selected from the group consisting of fragments comprising at least 10, at least 20, at least 30, at least 50 and more than 50 consecutive nucleotides of one of SEQ ID NOs: 8-3795.
- 6. The fragment of Paragraph 5, wherein said fragment is included in a nucleic acid obtained from an organism selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr
 (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes,
 Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria
- meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

7. The fragment of Paragraph 5, wherein said fragment is included in a nucleic acid obtained from an organism other than Escherichia coli.

8. A vector comprising a promoter operably linked to the nucleic acid of any one of Paragraphs 1-7.

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- 9. The vector of Paragraph 8, wherein said promoter is active in a microorganism selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefvr (also called Candida 10 pseudotropicalis). Candida dubliniensis, Chlamydia pneumoniae. Chlamydia trachomatus. Clostridium botulinum. Clostridium difficile. Clostridium perfringens. Coccidiodes immitis. Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae. Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri. Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.
 - 10. A host cell containing the vector of Paragraph 8 or Paragraph 9.
 - 11. A purified or isolated antisense nucleic acid comprising a nucleotide sequence complementary to at least a portion of an intragenic sequence, intergenic sequence, sequences spanning at least a portion of two or more genes, 5' noncoding region, or 3' noncoding region within an operon comprising a proliferation-required gene whose activity or expression is inhibited by an antisense nucleic acid comprising the nucleotide sequence of one of SEQ ID NOs.; 8-3795.
 - 12. The purified or isolated antisense nucleic acid of Paragraph 11, wherein said antisense nucleic acid is complementary to a nucleic acid from an organism selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corvnebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli,

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Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

- 13. The purified or isolated antisense nucleic acid of Paragraph 11, wherein said nucleotide sequence is complementary to a nucleotide sequence of a nucleic acid from an organism other than *E. coli*.
- 14. The purified or isolated antisense nucleic acid of Paragraph 11, wherein said proliferation-required gene comprises a nucleotide sequence selected from the group consisting of SEO ID NOS.: 3796-3800, 3806-4860, 5916-10012.
- 15. A purified or isolated nucleic acid comprising a nucleotide sequence having at least 70% identity to a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, fragments comprising at least 25 consecutive nucleotides of SEQ ID NOs.: 8-3795, the nucleotide sequences complementary to SEQ ID NOs.: 8-3795 and the sequences complementary to fragments comprising at least 25 consecutive nucleotides of SEQ ID NOs.: 8-3795 as determined using BLASTN version 2.0 with the default parameters.

16. The purified or isolated nucleic acid of Paragraph 15, wherein said nucleic acid is

obtained from an organism selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia. 25 Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata). Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, 30 Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinti, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, 35 Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus

pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

17. The nucleic acid of Paragraph 15, wherein said nucleic acid is obtained from an organism other than *E. coli*.

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- 18. A vector comprising a promoter operably linked to a nucleic acid encoding a polypeptide whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence of any one of SEQ ID NOs.: 8-3795.
- 19. The vector of Paragraph 18, wherein said nucleic acid encoding said polypeptide is obtained from an organism selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei. Candida kefvr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes. Moxarella catarrhalis. Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.
- 20. The vector of Paragraph 18, wherein said nucleotide sequence encoding said polypeptide is obtained from an organism other than *E. coli*.
 - 21. A host cell containing the vector of Paragraph 18.
- 22. The vector of Paragraph 18, wherein said polypeptide comprises a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NOs: 3801-3805, 4861-5915, 10013-14110.
 - 23. The vector of Paragraph 18, wherein said promoter is operably linked to a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012.
- 24. A purified or isolated polypeptide comprising a polypeptide whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence of any one of SEQ ID NOs.: 8-3795, or a fragment selected from the group consisting of fragments comprising at least 5,

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at least 10, at least 20, at least 30, at least 40, at least 50, at least 60 or more than 60 consecutive amino acids of one of the said polypeptides.

- 25. The polypeptide of Paragraph 24, wherein said polypeptide comprises an amino acid sequence of any one of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110 or a fragment comprising at least 5, at least 10, at least 20, at least 30, at least 40, at least 50, at least 60 or more than 60 consecutive amino acids of a polypeptide comprising an amino acid sequence selected from the group consisting of SEO ID NOs.: 3801-3805, 4861-5915, 10013-14110.
- 26. The polypeptide of Paragraph 24, wherein said polypeptide is obtained from an organism selected from the group consisting of Anaplasma marginale. Aspergillus fumigatus. Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, 10 Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes 15 immitis. Corvnebacterium diptheriae. Cryptococcus neoformans. Enterobacter cloacae. Enterococcus faecalis. Enterococcus faecium. Escherichia coli. Haemophilus influenzae. Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis 20 carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae. Shigella flexneri. Shigella sonnei. Staphylococcus epidermidis. Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis 25 and any species falling within the genera of any of the above species.
 - 27. The polypeptide of Paragraph 24, wherein said polypeptide is obtained from an organism other than *E. coli*.
 - 28. A purified or isolated polypeptide comprising a polypeptide having at least 25% amino acid identity to a polypeptide whose expression is inhibited by a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, or at least 25% amino acid identity to a fragment comprising at least 10, at least 20, at least 30, at least 40, at least 50, at least 60 or more than 60 consecutive amino acids of a polypeptide whose expression is inhibited by a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 as determined using FASTA version 3.0t78 with the default parameters.
 - 29. The polypeptide of Paragraph 28, wherein said polypeptide has at least 25% identity to a polypeptide comprising one of SEQ ID NOs: 3801-3805, 4861-5915, 10013-14110 or at least 25% identity to a fragment comprising at least 5, at least 10, at least 20, at least 30, at least 40, at

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least 50, at least 60 or more than 60 consecutive amino acids of a polypeptide comprising one of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110 as determined using FASTA version 3.0t78 with the default parameters.

- 30. The polypeptide of Paragraph 28, wherein said polypeptide is obtained from an organism selected from the group consisting of Anaplasma marginale, Aspergillus funigatus. Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia. Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae. Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis. Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.
- 31. The polypeptide of Paragraph 28, wherein said polypeptide is obtained from an organism other than *E. coli*.
- An antibody capable of specifically binding the polypeptide of one of Paragraphs
 28-31.
 - 33. A method of producing a polypeptide, comprising introducing a vector comprising a promoter operably linked to a nucleic acid comprising a nucleotide sequence encoding a polypeptide whose expression is inhibited by an antisense nucleic acid comprising one of SEQ ID NOs.: 8-3795 into a cell.
 - 34. The method of Paragraph 33, further comprising the step of isolating said polypeptide.
 - 35. The method of Paragraph 33, wherein said polypeptide comprises an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
- 36. The method of Paragraph 33, wherein said nucleic acid encoding said polypeptide is obtained from an organism selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata).

Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefvr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corvnebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae. 5 Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemonhilus influenzae Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis. 10 Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

37. The method of Paragraph 33, wherein said nucleic acid encoding said polypeptide is obtained from an organism other than *E. coli*.

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38. The method of Paragraph 33, wherein said promoter is operably linked to a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012.

39. A method of inhibiting proliferation of a cell in an individual comprising inhibiting the activity or reducing the amount of a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 or inhibiting the activity or reducing the amount of a nucleic acid encoding said gene product.

40. The method of Paragraph 39, wherein said method comprises inhibiting said activity or reducing said amount of a gene product in an organism selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus yulgaris, Pseudomonas aeruginosa, Salmonella bongori,

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Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

- 41. The method of Paragraph 39, wherein said method comprises inhibiting said activity or reducing said amount of a gene product in an organism other than E. coli.
- 42. The method of Paragraph 39, wherein said gene product is present in an organism other than *E. coli*.
- 43. The method of Paragraph 39, wherein said gene product comprises a polypeptide comprising a sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
- 44. A method for identifying a compound which influences the activity of a gene product required for proliferation, said gene product comprising a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, said method comprising:

contacting said gene product with a candidate compound; and determining whether said compound influences the activity of said gene product.

45. The method of Paragraph 44, wherein said gene product is from an organism selected 20 from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus. 25 Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis. Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, 30 Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, 35 Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

46. The method of Paragraph 44, wherein said gene product is from an organism other than *E. coli*.

- 47. The method of Paragraph 44, wherein said gene product is a polypeptide and said activity is an enzymatic activity.
- 48. The method of Paragraph 44, wherein said gene product is a polypeptide and said activity is a carbon compound catabolism activity.

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- 49. The method of Paragraph 44, wherein said gene product is a polypeptide and said activity is a biosynthetic activity.
- 50. The method of Paragraph 44, wherein said gene product is a polypeptide and said activity is a transporter activity.
 - 51. The method of Paragraph 44, wherein said gene product is a polypeptide and said activity is a transcriptional activity.
- 52. The method of Paragraph 44, wherein said gene product is a polypeptide and said activity is a DNA replication activity.
- 53. The method of Paragraph 44, wherein said gene product is a polypeptide and said activity is a cell division activity.
 - 54. The method of Paragraph 44, wherein said gene product is an RNA.
- 55. The method of Paragraph 44, wherein said gene product is a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
 - 56. A compound identified using the method of Paragraph 44.
- 57. A method for identifying a compound or nucleic acid having the ability to reduce the activity or level of a gene product required for proliferation, said gene product comprising a gene product whose activity or expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, said method comprising:
 - (a) contacting a target gene or RNA encoding said gene product with a candidate compound or nucleic acid; and
 - (b) measuring an activity of said target.
- 58. The method of Paragraph 57, wherein said target gene or RNA is from an organism selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus

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faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori,

Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

- 59. The method of Paragraph 57, wherein said target gene or RNA is from an organism other than *E. coli*.
- 60. The method of Paragraph 57, wherein said gene product is from an organism other than *E. coli*.
- 15 61. The method of Paragraph 57, wherein said target is a messenger RNA molecule and said activity is translation of said messenger RNA.
 - 62. The method of Paragraph 57, wherein said target is a messenger RNA molecule and said activity is transcription of a gene encoding said messenger RNA.
 - 63. The method of Paragraph 57, wherein said target is a gene and said activity is transcription of said gene.
 - 64. The method of Paragraph 57, wherein said target is a nontranslated RNA and said activity is processing or folding of said nontranslated RNA or assembly of said nontranslated RNA into a protein/RNA complex.
 - 65. The method of Paragraph 57, wherein said target is a messenger RNA molecule encoding a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
 - 66. The method of Paragraph 57, wherein said target comprises a nucleic acid selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012.
 - 67. A compound or nucleic acid identified using the method of Paragraph 57.
 - 68. A method for identifying a compound which reduces the activity or level of a gene product required for proliferation of a cell, wherein the activity or expression of said gene product is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, said method comprising the steps of:
 - (a) providing a sublethal level of an antisense nucleic acid comprising a nucleotide sequence complementary to a nucleic acid comprising a nucleotide sequence encoding said gene product in a cell to reduce the activity or amount of said gene product in said cell, thereby producing a sensitized cell;

- (b) contacting said sensitized cell with a compound; and
- (c) determining the degree to which said compound inhibits proliferation of said sensitized cell relative to a cell which does not contain said antisense nucleic acid.
- 69. The method of Paragraph 68, wherein said determining step comprises determining whether said compound inhibits the growth of said sensitized cell to a greater extent than said compound inhibits the growth of a nonsensitized cell.

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- 70. The method of Paragraph 68, wherein said cell is a Gram positive bacterium.
- 71. The method of Paragraph 68, wherein said Gram positive bacterium is selected from the group consisting of *Staphylococcus* species, *Streptococcus* species, *Enterococcus* species, *Mycobacterium* species, *Clostridium* species, and *Bacillus* species.
 - 72. The method of Paragraph 68, wherein said bacterium is Staphylococcus aureus.
- 73. The method of Paragraph 72, wherein said *Staphylococcus* species is coagulase negative.
- 74. The method of Paragraph 72, wherein said bacterium is selected from the group consisting of *Staphylococcus aureus* RN450 and *Staphylococcus aureus* RN4220.
- 75. The method of Paragraph 68, wherein said cell is an organism selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida 20 guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, 25 Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella 30 boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis. Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica.
 - Yersinia pestis and any species falling within the genera of any of the above species.

 76. The method of Paragraph 68, wherein said cell is not an E. coli cell.
- 77. The method of Paragraph 68, wherein said gene product is from an organism other than 35 E. coli.
 - 78. The method of Paragraph 68, wherein said antisense nucleic acid is transcribed from an inducible promoter.

79. The method of Paragraph 68, further comprising the step of contacting said cell with a concentration of inducer which induces transcription of said antisense nucleic acid to a sublethal level.

- 80. The method of Paragraph 68, wherein growth inhibition is measured by monitoring optical density of a culture growth solution.
 - 81. The method of Paragraph 68, wherein said gene product is a polypeptide.
 - 82. The method of Paragraph 81, wherein said polypeptide comprises an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
 - 83. The method of Paragraph 68, wherein said gene product is an RNA.
- 84. The method of Paragraph 68, wherein nucleic acid encoding said gene product comprises a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012.
 - 85. A compound identified using the method of Paragraph 68.

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- 86. A method for inhibiting cellular proliferation comprising introducing an effective amount of a compound with activity against a gene whose activity or expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 or a compound with activity against the product of said gene into a population of cells expressing said gene.
- . 87. The method of Paragraph 86, wherein said compound is an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, or a proliferation-inhibiting portion thereof.
- 88. The method of Paragraph 86, wherein said proliferation inhibiting portion of one of SEQ ID NOs.: 8-3795 is a fragment comprising at least 10, at least 20, at least 25, at least 30, at least 50 or more than 51 consecutive nucleotides of one of SEO ID NOs.: 8-3795.
- 89. The method of Paragraph 86, wherein said population is a population of Gram positive bacteria.
- 90. The method of Paragraph 89, wherein said population of Gram positive bacteria is selected from the group consisting of a population of *Staphylococcus* species, *Streptococcus* species, *Enterococcus* species, *Mycobacterium* species, *Clostridium* species, and *Bacillus* species.
- 91. The method of Paragraph 86, wherein said population is a population of Staphylococcus aureus.
- 92. The method of Paragraph 91, wherein said population is a population of a bacterium selected from the group consisting of *Staphylococcus aureus* RN450 and *Staphylococcus aureus* RN4220.
- 93. The method of Paragraph 86, wherein said population is a population of a bacterium selected from the group consisting of *Anaplasma marginale*, *Aspergillus fumigatus*, *Bacillus*

anthracis. Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis. Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, 5 Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum. Klebsiella pneumoniae. Listeria monocytogenes, Mycobacterium leprae. Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, 10 Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans. 15 Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

- 94. The method of Paragraph 86, wherein said population is a population of an organism other than E. coli.
- 95. The method of Paragraph 86, wherein said product of said gene is from an organism other than *E. coli*.

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- 96. The method of Paragraph 86, wherein said gene encodes a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
- 97. The method of Paragraph 86, wherein said gene comprises a nucleotide sequence selected from the group consisting of SEO ID NOS.: 3796-3800, 3806-4860, 5916-10012.
 - 98. A composition comprising an effective concentration of an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, or a proliferation-inhibiting portion thereof in a pharmaceutically acceptable carrier.
 - 99. The composition of Paragraph 98, wherein said proliferation-inhibiting portion of one of SEQ ID NOs.: 8-3795 comprises at least 20, at least 25, at least 30, at least 50 or more than 50 consecutive nucleotides of one of SEQ ID NOs.: 8-3795.
 - 100. A method for inhibiting the activity or expression of a gene in an operon required for proliferation wherein the activity or expression of at least one gene in said operon is inhibited by an antisense nucleic acid comprising a sequence selected from the group consisting of SEQ ID NOs.: 8-3795, said method comprising contacting a cell in a cell population with an antisense nucleic acid complementary to at least a portion of said operon.

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101. The method of Paragraph 100, wherein said antisense nucleic acid comprises a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 or a proliferation-inhibiting portion thereof.

- 102. The method of Paragraph 100, wherein said cell is selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata). Candida tropicalis. Candida parapsilosis. Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corvnebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae. Enterococcus faecalis. Enterococcus faecium. Escherichia coli Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica. Yersinia pestis and any species falling within the genera of any of the above species.
 - 103. The method of Paragraph 100, wherein said cell is not an E. coli cell.
- 104. The method of Paragraph 100, wherein said gene is from an organism other than E. coli.
- 105. The method of Paragraph 100, wherein said cell is contacted with said antisense nucleic acid by introducing a plasmid which expresses said antisense nucleic acid into said cell population.
 - 106. The method of Paragraph 100, wherein said cell is contacted with said antisense nucleic acid by introducing a phage which encodes said antisense nucleic acid into said cell population.
 - 107. The method of Paragraph 100, wherein said cell is contacted with said antisense nucleic acid by expressing said antisense nucleic acid from the chromosome of cells in said cell population.
 - 108. The method of Paragraph 100, wherein said cell is contacted with said antisense nucleic acid by introducing a promoter adjacent to a chromosomal copy of said antisense nucleic acid such that said promoter directs the transcription of said antisense nucleic acid.

109. The method of Paragraph 100, wherein said cell is contacted with said antisense nucleic acid by introducing a retron which expresses said antisense nucleic acid into said cell population.

110. The method of Paragraph 100, wherein said cell is contacted with said antisense nucleic acid by introducing a ribozyme into said cell-population, wherein a binding portion of said ribozyme comprises said antisense nucleic acid.

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- 111. The method of Paragraph 100, wherein said cell is contacted with said antisense nucleic acid by introducing a liposome comprising said antisense nucleic acid into said cell.
- 112. The method of Paragraph 100, wherein said cell is contacted with said antisense nucleic acid by electroporation of said antisense nucleic acid into said cell.
- 113. The method of Paragraph 100, wherein said antisense nucleic acid is a fragment comprising at least 10, at least 20, at least 25, at least 30, at least 50 or more than 50 consecutive nucleotides of one of SEQ ID NOs.: 8-3795.
- 114. The method of Paragraph 100 wherein said antisense nucleic acid is a synthetic oligonucleotide.
- 115. The method of Paragraph 100, wherein said gene comprises a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012.
- 116. A method for identifying a gene which is required for proliferation of a cell comprising:
 - (a) contacting a cell with an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, wherein said cell is a cell other than the organism from which said nucleic acid was obtained;
 - (b) determining whether said nucleic acid inhibits proliferation of said cell; and
 - (c) identifying the gene in said cell which encodes the mRNA which is complementary to said antisense nucleic acid or a portion thereof.
- 117. The method of Paragraph 116, wherein said cell is selected from the group consisting of *Staphylococcus* species, *Streptococcus* species, *Enterococcus* species, *Mycobacterium* species, *Clostridium* species, and *Bacillus* species.
- 118. The method of Paragraph 116 wherein said cell is selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae,

Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

119. The method of Paragraph 116, wherein said cell is not E. coli.

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- 120. The method of Paragraph 116, further comprising operably linking said antisense nucleic acid to a promoter which is functional in said cell, said promoter being included in a vector, and introducing said vector into said cell.
- 121. A method for identifying a compound having the ability to inhibit proliferation of a cell comprising:
 - (a) identifying a homolog of a gene or gene product whose activity or level is inhibited by a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs. 8-3795 in a test cell, wherein said test cell is not the cell from which said nucleic acid was obtained;
 - (b) identifying an inhibitory nucleic acid sequence which inhibits the activity of said homolog in said test cell;
 - (c) contacting said test cell with a sublethal level of said inhibitory nucleic acid, thus sensitizing said cell;
 - (d) contacting the sensitized cell of step (c) with a compound; and
 - (e) determining the degree to which said compound inhibits proliferation of said sensitized cell relative to a cell which does not contain said inhibitory nucleic acid.
- 122. The method of Paragraph 121, wherein said determining step comprises determining whether said compound inhibits proliferation of said sensitized test cell to a greater extent than said compound inhibits proliferation of a nonsensitized test cell.
- 123. The method of Paragraph 121, wherein step (a) comprises identifying a nucleic acid homologous to a gene or gene product whose activity or level is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs. 8-3795 or a nucleic acid encoding a homologous polypeptide to a polypeptide whose activity or level is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs. 8-3795 by using an algorithm selected from the group consisting of BLASTN version 2.0 with the default parameters and FASTA version 3.0t78 algorithm with the default parameters to identify said homologous nucleic acid or said nucleic acid encoding a homologous polypeptide in a database.

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124. The method of Paragraph 121 wherein said step (a) comprises identifying a homologous nucleic acid or a nucleic acid comprising a sequence of nucleotides encoding a homologous polypeptide by identifying nucleic acids which hybridize to said nucleic acid selected from the group consisting of SEQ ID NOs. 8-3795 or the complement of said nucleic acid selected from the group consisting of SEQ ID NOs. 8-3795.

- 125. The method of Paragraph 121 wherein step (a) comprises expressing a nucleic acid selected from the group consisting of SEQ ID NOs. 8-3795 in said test cell.
- 126. The method of Paragraph 121, wherein step (a) comprises identifying a homologous nucleic acid or a nucleic acid encoding a homologous polypeptide in a test cell 10 selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter ieiuni Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis. Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, 15 Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis. Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae. Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides. 20 Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, 25 Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.
 - 127. The method of Paragraph 121, wherein step (a) comprises identifying a homologous nucleic acid or a nucleic acid encoding a homologous polypeptide in a test cell other than *E. coli*.
 - 128. The method of Paragraph 121, wherein said inhibitory nucleic acid is an antisense nucleic acid.
 - 129. The method of Paragraph 121, wherein said inhibitory nucleic acid comprises an antisense nucleic acid to a portion of said homolog.
- 130. The method of Paragraph 121, wherein said inhibitory nucleic acid comprises anantisense nucleic acid to a portion of the operon encoding said homolog.

131. The method of Paragraph 121, wherein the step of contacting the cell with a sublethal level of said inhibitory nucleic acid comprises directly contacting the surface of said cell with said inhibitory nucleic acid.

132. The method of Paragraph 121, wherein the step of contacting the cell with a sublethal level of said inhibitory nucleic acid comprises transcribing an antisense nucleic acid complementary to at least a portion of the RNA transcribed from said homolog in said cell.

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- 133. The method of Paragraph 121, wherein said gene product comprises a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
- 134. The method of Paragraph 121, wherein said gene comprises a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012.
 - 135. A compound identified using the method of Paragraph 121.
- 136. A method of identifying a compound having the ability to inhibit proliferation comprising:
 - (a) contacting a test cell with a sublethal level of a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs. 8-3795 or a portion thereof which inhibits the proliferation of the cell from which said nucleic acid was obtained, thus sensitizing said test cell;
 - (b) contacting the sensitized test cell of step (a) with a compound; and
 - (c) determining the degree to which said compound inhibits proliferation of said sensitized test cell relative to a cell which does not contain said nucleic acid.
- 137. The method of Paragraph 136, wherein said determining step comprises determining whether said compound inhibits proliferation of said sensitized test cell to a greater extent than said compound inhibits proliferation of a nonsensitized test cell.
 - 138. A compound identified using the method of Paragraph 136.
- 139. The method of Paragraph 136, wherein said test cell is selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori,

Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

140. The method of Paragraph 136, wherein the test cell is not E. coli.

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- 141. A method for identifying a compound having activity against a biological pathway required for proliferation comprising:
 - (a) sensitizing a cell by providing a sublethal level of an antisense nucleic acid complementary to a nucleic acid encoding a gene product required for proliferation, wherein the activity or expression of said gene product is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, in said cell to reduce the activity or amount of said gene product;
 - (b) contacting the sensitized cell with a compound; and
 - (c) determining the degree to which said compound inhibits the growth of said sensitized cell relative to a cell which does not contain said antisense nucleic acid.
- 142. The method of Paragraph 141, wherein said determining step comprises determining whether said compound inhibits the growth of said sensitized cell to a greater extent than said compound inhibits the growth of a nonsensitized cell.
- 143. The method of Paragraph 141, wherein said cell is selected from the group consisting of bacterial cells, fungal cells, plant cells, and animal cells.
 - 144. The method of Paragraph 141, wherein said cell is a Gram positive bacterium.
- 145. The method of Paragraph 144, wherein said Gram positive bacterium is selected from the group consisting of *Staphylococcus* species, *Streptococcus* species, *Enterococcus* species, *Mycobacterium* species, *Clostridium* species, and *Bacillus* species.
- 146. The method of Paragraph 145, wherein said Gram positive bacterium is *Staphylococcus aureus*.
- 147. The method of Paragraph 146, wherein said Gram positive bacterium is selected from the group consisting of Staphylococcus aureus RN450 and Staphylococcus aureus RN4220.
- 148. The method of Paragraph 141, wherein said cell is selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli,

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Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

149. The method of Paragraph 141, wherein said cell is not an E. coli cell.

- 150. The method of Paragraph 141, wherein said gene product is from an organism other than *E. coli*.
- 151. The method of Paragraph 141, wherein said antisense nucleic acid is transcribed from an inducible promoter.
- 152. The method of Paragraph 141, further comprising contacting the cell with an agent which induces transcription of said antisense nucleic acid from said inducible promoter, wherein said antisense nucleic acid is transcribed at a sublethal level.
- 153. The method of Paragraph 141, wherein inhibition of proliferation is measured by monitoring the optical density of a liquid culture.
- 154. The method of Paragraph 141, wherein said gene product comprises a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
- 155. The method of Paragraph 141, wherein said nucleic acid encoding said gene product comprises a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012.
 - 156. A compound identified using the method of Paragraph 141.
- 157. A method for identifying a compound having the ability to inhibit cellular proliferation comprising:
 - (a) contacting a cell with an agent which reduces the activity or level of a gene product required for proliferation of said cell, wherein said gene product is a gene product whose activity or expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795;
 - (b) contacting said cell with a compound; and
 - (c) determining whether said compound reduces proliferation of said contacted cell by acting on said gene product.

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158. The method of Paragraph 157, wherein said determining step comprises determining whether said compound reduces proliferation of said contacted cell to a greater extent than said compound reduces proliferation of cells which have not been contacted with said agent.

- The method of Paragraph 157, wherein said cell is selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis. Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis. Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori. Salmonella cholerasuis, Salmonella enterica, Salmonella varatyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.
 - 160. The method of Paragraph 157, wherein said cell is not an E. coli cell,
- 161. The method of Paragraph 157, wherein said gene product is from an organism other than *E. coli*.
- 162. The method of Paragraph 157, wherein said agent which reduces the activity or level of a gene product required for proliferation of said cell comprises an antisense nucleic acid to a gene or operon required for proliferation.
- 163. The method of Paragraph 157, wherein said agent which reduces the activity or level of a gene product required for proliferation of said cell comprises a compound known to inhibit growth or proliferation of a cell.
- 164. The method of Paragraph 157, wherein said cell contains a mutation which reduces the activity or level of said gene product required for proliferation of said cell.
- 165. The method of Paragraph 157, wherein said mutation is a temperature sensitive mutation.
- 166. The method of Paragraph 157, wherein said gene product comprises a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
 - 167. A compound identified using the method of Paragraph 157.

168. A method for identifying the biological pathway in which a proliferation-required gene or its gene product lies, wherein said gene or gene product comprises a gene or gene product whose activity or expression is inhibited by an antisense nucleic acid comprising a sequence selected from the group consisting of SEQ ID NOs.: 8-3795, said method comprising:

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- (a) providing a sublethal level of an antisense nucleic acid which inhibits the activity of said proliferation-required gene or gene product in a test cell;
- (b) contacting said test cell with a compound known to inhibit growth or proliferation of a cell, wherein the biological pathway on which said compound acts is known; and

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- (c) determining the degree to which said proliferation of said test cell is inhibited relative to a cell which was not contacted with said compound.
- 169. The method of Paragraph 168, wherein said determining step comprises determining whether said test cell has a substantially greater sensitivity to said compound than a cell which does not express said sublethal level of said antisense nucleic acid.
- 170. The method of Paragraph 168, wherein said gene product comprises a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
- 171. The method of Paragraph 168, wherein said test cell is selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.
 - 172. The method of Paragraph 168, wherein said test cell is not an E. coli cell.
 - 173. The method of Paragraph 168, wherein said gene product is from an organism other than *E. coli*.

174. A method for determining the biological pathway on which a test compound acts comprising:

- (a) providing a sublethal level of an antisense nucleic acid complementary to a proliferation-required nucleic acid in a first cell, wherein the activity or expression of said proliferation-required nucleic acid is inhibited by an antisense nucleic acid comprising a sequence selected from the group consisting of SEQ ID NOs.: 8-3795 and wherein the biological pathway in which said proliferation-required nucleic acid or a protein encoded by said proliferation-required nucleic acid lies is known,
 - (b) contacting said first cell with said test compound; and
- (c) determining the degree to which said test compound inhibits proliferation of said first cell relative to a cell which does not contain said antisense nucleic acid.
- 175. The method of Paragraph 174, wherein said determining step comprises determining whether said first cell has a substantially greater sensitivity to said test compound than a cell which does not express said sublethal level of said antisense nucleic acid.
 - 176. The method of Paragraph 174, further comprising:

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- (d) providing a sublethal level of a second antisense nucleic acid complementary to a second proliferation-required nucleic acid in a second cell, wherein said second proliferation-required nucleic acid is in a different biological pathway than said proliferation-required nucleic acid in step (a); and
- (e) determining whether said second cell does not have a substantially greater sensitivity to said test compound than a cell which does not express said sublethal level of said second antisense nucleic acid, wherein said test compound is specific for the biological pathway against which the antisense nucleic acid of step (a) acts if said first cell has a substantially greater sensitivity to said test compound than said second cell.
- 177. The method of Paragraph 174, wherein said first cell is selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella

typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

178. The method of Paragraph 174, wherein said first cell is not an E. coli cell.

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- 179. The method of Paragraph 174, wherein said proliferation-required nucleic acid is from an organism other than *E. coli*.
- 180. A purified or isolated nucleic acid comprising a sequence selected from the group consisting of SEQ ID NOs.: 8-3795.
- 181. A compound which interacts with a gene or gene product whose activity or expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence of one of SEQ ID NOs.: 8-3795 to inhibit proliferation.
- 182. The compound of Paragraph 181, wherein said gene product is a polypeptide comprising one of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
- 183. The compound of Paragraph 181, wherein said gene comprises a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012.
- 184. A compound which interacts with a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence of one of SEQ ID NOs.: 8-3795 to inhibit proliferation.
- 185. A method for manufacturing an antibiotic comprising the steps of:
 screening one or more candidate compounds to identify a compound that reduces the
 activity or level of a gene product required for proliferation, said gene product comprising a gene
 product whose activity or expression is inhibited by an antisense nucleic acid comprising a
 nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795; and
- 186. The method of Paragraph 185, wherein said screening step comprises performing any one of the methods of Paragraphs 44, 68, 121, 136, 141, and 157.
- 187. The method of Paragraph 185, wherein said gene product is a polypeptide comprising one of SEQ ID NOs:3801-3805, 4861-5915, 10013-14110.

manufacturing the compound so identified.

- 188. A method for inhibiting proliferation of a cell in a subject comprising administering an effective amount of a compound that reduces the activity or level of a gene product required for proliferation of said cell, said gene product comprising a gene product whose activity or expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 to said subject.
- 189. The method of Paragraph 188 wherein said subject is selected from the group consisting of vertebrates, mammals, avians, and human beings.

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190. The method of Paragraph 188, wherein said gene product comprises a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.

- 191. The method of Paragraph 188, wherein said cell is selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata). Candida tropicalis. Candida parapsilosis. Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis). Candida dubliniensis, Chlamydia pneumoniae. Chlamydia trachomatus. Clostridium hotulinum. Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans. Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae. Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella hongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica. Yersinia pestis and any species falling within the genera of any of the above species.
 - 192. The method of Paragraph 188, wherein said cell is not E. coli.
- 193. The method of Paragraph 188, wherein said gene product is from an organism other than *E. coli*.
- 194. A purified or isolated nucleic acid consisting essentially of the coding sequence of one of SEQ ID NOs: 3796-3800, 3806-4860, 5916-10012.
 - 195. A fragment of the nucleic acid of Paragraph 8, said fragment comprising at least 10, at least 20, at least 25, at least 30, at least 50 or more than 50 consecutive nucleotides of one of SEQ ID NOs: 3796-3800, 3806-4860, 5916-10012.
 - 196. A purified or isolated nucleic acid comprising a nucleic acid having at least 70% nucleotide sequence identity to a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, fragments comprising at least 25 consecutive nucleotides of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, the nucleotide sequences complementary to SEQ ID NOs.:3796-3800, 3806-4860, 5916-10012, and the nucleotide sequences complementary to fragments comprising at least 25 consecutive nucleotides of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012 as determined using BLASTN version 2.0 with the default parameters.

The nucleic acid of Paragraph 196, wherein said nucleic acid is from an organism 197. selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida 5 pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis. Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, 10 Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae. Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides. Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri. 15 Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans. Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

198. The nucleic acid of Paragraph 196, wherein said nucleic acid is from an organism other than *E. coli*.

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A method of inhibiting proliferation of a cell comprising inhibiting the activity or 199. reducing the amount of a gene product in said cell or inhibiting the activity or reducing the amount of a nucleic acid encoding said gene product in said cell, wherein said gene product is selected from the group consisting of a gene product having having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, a gene product encoded by a nucleic acid which hybridizes to a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid which hybridizes to a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795

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under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs: 8-3795.

- The method of Paragraph 199, wherein said method comprises inhibiting said 200. activity or reducing said amount of said gene product or inhibiting the activity or reducing the amount of a nucleic acid encoding said gene product in an organism selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia. Campylobacter jejuni. Candida albicans. Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis). Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis. Corvnebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli. Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum. Klebsiella pneumoniae. Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica. Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella hongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis. Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.
- 201. The method of Paragraph 199, wherein said method comprises inhibiting said activity or reducing said amount of said gene product or inhibiting the activity or reducing the amount of a nucleic acid encoding said gene product in an organism other than *E. coli*.
- 202. The method of Paragraph 199, wherein said gene product is from an organism other than *E. coli*.
- 203. The method of Paragraph 199, wherein said gene product comprises a polypeptide selected from the group consisting of a polypeptide having at least 25% amino acid identity as determined using FASTA version 3.0t78 to a polypeptide selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110 and a polypeptide whose activity may be complemented by a polypeptide selected from the group consisting of SEQ ID NOs: 3801-3805, 4861-5915, 10013-14110.
- 204. The method of Paragraph 199, wherein said gene product is encoded by a nucleic acid selected from the group consisting of a nucleic acid comprising a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-

3800, 3806-4860, 5916-10012, a nucleic acid comprising a nucleotide sequence which hybridizes to a sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800. 3806-4860. 5916-10012 under moderate conditions.

205. A method for identifying a compound which influences the activity of a gene product required for proliferation comprising:

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contacting a candidate compound with a gene product selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795; and

determining whether said candidate compound influences the activity of said gene product.

206. The method of Paragraph 205, wherein said gene product is from an organism selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus

faecalis. Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori,

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Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

- 207. The method of Paragraph 205, wherein said gene product is from an organism other than *E. coli*.
- 208. The method of Paragraph 205, wherein said gene product is a polypeptide selected from the group consisting of a polypeptide having at least 25% amino acid identity as determined using FASTA version 3.0t78 to a polypeptide selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110 and a polypeptide whose activity may be complemented by a polypeptide selected from the group consisting of SEQ ID NOs: 3801-3805, 4861-5915, 10013-14110.
- 209. The method of Paragraph 205, wherein said gene product is encoded by a nucleic acid selected from the group consisting of a nucleic acid comprising a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, a nucleic acid which hybridizes to a sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under stringent conditions, and a nucleic acid which hybridizes to a sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under moderate conditions.
 - 210. A compound identified using the method of Paragraph 205.
- 211. A method for identifying a compound or nucleic acid having the ability to reduce the activity or level of a gene product required for proliferation comprising:
 - (a) providing a target that is a gene or RNA, wherein said target comprises a nucleic acid that encodes a gene product selected from the group consisting of a gene product having having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleic acid identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group

consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795:

- (b) contacting said target with a candidate compound or nucleic acid; and
- (c) measuring an activity of said target.

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- 212. The method of Paragraph 211, wherein said target gene or RNA is from an 15 organism selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis. Bacterioides fravilis Bordetella pertussis. Burkholderia cepacia. Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata). Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia 20 trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae. Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria 25 meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus 30 pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.
 - 213. The method of Paragraph 211, wherein said target gene or RNA is from an organism other than *E. coli*.
- 214. The method of Paragraph 211, wherein said gene product is from an organism other than E. coli.
 - 215. The method of Paragraph 211, wherein said target is a messenger RNA molecule and said activity is translation of said messenger RNA.

216. The method of Paragraph 211, wherein said compound is a nucleic acid and said activity is translation of said gene product.

- 217. The method of Paragraph 211, wherein said target is a gene and said activity is transcription of said gene.
- 218. The method of Paragraph 211, wherein said target is a nontranslated RNA and said activity is processing or folding of said nontranslated RNA or assembly of said nontranslated RNA into a protein/RNA complex.
- 219. The method of Paragraph 211, wherein said target gene is a messenger RNA molecule encoding a polypeptide selected from the group consisting of a polypeptide having at least 25% amino acid identity as determined using FASTA version 3.0t78 to a polypeptide selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110 and a polypeptide whose activity may be complemented by a polypeptide selected from the group consisting of SEQ ID NOs: 3801-3805, 4861-5915, 10013-14110.
- 220. The method of Paragraph 11, wherein said target gene comprises a nucleic acid selected from the group consisting of a nucleic acid comprising a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, a nucleic acid which hybridizes to a sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under stringent conditions, and a nucleic acid which hybridizes to a sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under moderate conditions.
 - 221. A compound or nucleic acid identified using the method of Paragraph 211.
- 222. A method for identifying a compound which reduces the activity or level of a gene product required for proliferation of a cell comprising:

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(a) providing a sublethal level of an antisense nucleic acid complementary to a nucleic acid encoding said gene product in a cell to reduce the activity or amount of said gene product in said cell, thereby producing a sensitized cell, wherein said gene product is selected from the group consisting of a gene product having having at least 70% nucleic acid identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited

by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEO ID NOs: 8-3795:

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- (b) contacting said sensitized cell with a compound; and
- (c) determining the degree to which said compound inhibits the growth of said sensitized cell relative to a cell which does not contain said antisense nucleic acid.
- 223. The method of Paragraph 222, wherein said determining step comprises determining whether said compound inhibits the growth of said sensitized cell to a greater extent than said compound inhibits the growth of a nonsensitized cell.
- 224. The method of Paragraph 222, wherein said sensitized cell is a Gram positive bacterium.
- 225. The method of Paragraph 224, wherein said Gram positive bacterium is selected from the group consisting of *Staphylococcus* species, *Streptococcus* species, *Enterococcus* species, *Mycobacterium* species, *Clostridium* species, and *Bacillus* species.
 - 226. The method of Paragraph 225, wherein said bacterium is Staphylococcus aureus.
- 227. The method of Paragraph 224, wherein said *Staphylococcus* species is coagulase negative.
- 228. The method of Paragraph 226, wherein said bacterium is selected from the group consisting of *Staphylococcus aureus* RN450 and *Staphylococcus aureus* RN4220.
 - 229. The method of Paragraph 222, wherein said sensitized cell is an organism selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris,

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Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

- 230. The method of Paragraph 222, wherein said cell is an organism other than E. coli.
- 231. The method of Paragraph 222, wherein said gene product is from an organism other than *E. coli*.
- 232. The method of Paragraph 222, wherein said antisense nucleic acid is transcribed from an inducible promoter.
 - 233. The method of Paragraph 222, further comprising the step of contacting said cell with a concentration of inducer which induces transcription of said antisense nucleic acid to a sublethal level.
- 234. The method of Paragraph 222, wherein growth inhibition is measured by monitoring optical density of a culture medium.
 - 235. The method of Paragraph 222, wherein said gene product is a polypeptide.
- 236. The method of Paragraph 235, wherein said polypeptide comprises a polypeptide selected from the group consisting of a polypeptide having at least 25% amino acid identity as determined using FASTA version 3.0t78 to a polypeptide selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110 and a polypeptide whose activity may be complemented by a polypeptide selected from the group consisting of SEQ ID NOs: 3801-3805, 4861-5915, 10013-14110.
 - 237. The method of Paragraph 222, wherein said gene product is an RNA.
- 238. The method of Paragraph 222, wherein said nucleic acid encoding said gene product comprises a nucleic acid selected from the group consisting of a nucleic acid comprising a nucleic acid having at least 70% nucleic acid identity as determined using BLASTN version 2.0 with the default parameters to a sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, a nucleic acid which hybridizes to a sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under stringent conditions, and a nucleic acid which hybridizes to a sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under moderate conditions.
 - 239. A compound identified using the method of Paragraph 222.
- 240. A method for inhibiting cellular proliferation comprising introducing a compound with activity against a gene product or a compound with activity against a gene encoding said gene product into a population of cells expressing said gene product, wherein said gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence

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identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.; 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.; 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795.

- 241. The method of Paragraph 240, wherein said compound is an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, or a proliferation-inhibiting portion thereof.
- 20 242. The method of Paragraph 240, wherein said proliferation inhibiting portion of one of SEQ ID NOs.: 8-3795 is a fragment comprising at least 10, at least 20, at least 25, at least 30, at least 50 or more than 51 consecutive nucleotides of one of SEQ ID NOs.: 8-3795.
 - 243. The method of Paragraph 240, wherein said population is a population of Gram positive bacteria.
 - 244. The method of Paragraph 243, wherein said population of Gram positive bacteria is selected from the group consisting of a population of *Staphylococcus* species, *Streptococcus* species, *Enterococcus* species, *Mycobacterium* species, *Clostridium* species, and *Bacillus* species.
 - 245. The method of Paragraph 243, wherein said population is a population of Staphylococcus aureus.
 - 246. The method of Paragraph 245, wherein said population is a population of a bacterium selected from the group consisting of *Staphylococcus aureus* RN450 and *Staphylococcus aureus* RN4220.
 - 247. The method of Paragraph 240, wherein said population is a population of a bacterium selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei. Candida kefur

(also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae,
 Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium,
 Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

248. The method of Paragraph 240, wherein said population is a population of an organism other than *E. coli*.

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- 249. The method of Paragraph 240, wherein said product of said gene is from an organism other than *E. coli*.
- 250. The method of Paragraph 240, wherein said gene product is selected from the group consisting of a polypeptide having at least 25% amino acid identity as determined using FASTA version 3.0t78 to a polypeptide selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110 and a polypeptide whose activity may be complemented by a polypeptide selected from the group consisting of SEQ ID NOs: 3801-3805, 4861-5915, 10013-14110.
- 251. The method of Paragraph 240, wherein said gene comprises a nucleic acid selected from the group consisting of a nucleic acid comprising a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under moderate conditions.
 - 252. A preparation comprising an effective concentration of an antisense nucleic acid in a pharmaceutically acceptable carrier wherein said antisense nucleic acid is selected from the group consisting of a nucleic acid comprising a sequence having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 or a proliferation-inhibiting portion

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increol, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions.

- 253. The preparation of Paragraph 252, wherein said proliferation-inhibiting portion of one of SEQ ID NOs.: 8-3795 comprises at least 10, at least 20, at least 25, at least 30, at least 50 or more than 50 consecutive nucleotides of one of SEQ ID NOs.: 8-3795.
- 254. A method for inhibiting the activity or expression of a gene in an operon which encodes a gene product required for proliferation comprising contacting a cell in a cell population with an antisense nucleic acid comprising at least a proliferation-inhibiting portion of said operon in an antisense orientation, wherein said gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795. a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795.
- 255. The method of Paragraph 254, wherein said antisense nucleic acid comprises a nucleotide sequence having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a proliferation inhibiting portion thereof, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, and a nucleic acid which comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions.
- 256. The method of Paragraph 254, wherein said cell is selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis

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Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhin, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

- 257. The method of Paragraph 254, wherein said cell is not an E. coli cell.
- 258. The method of Paragraph 254, wherein said gene is from an organism other than E. coli.
- 259. The method of Paragraph 254, wherein said cell is contacted with said antisense nucleic acid by introducing a plasmid which transcribes said antisense nucleic acid into said cell population.
 - 260. The method of Paragraph 254, wherein said cell is contacted with said antisense nucleic acid by introducing a phage which transcribes said antisense nucleic acid into said cell population.
 - 261. The method of Paragraph 254, wherein said cell is contacted with said antisense nucleic acid by transcribing said antisense nucleic acid from the chromosome of cells in said cell population.
 - 262. The method of Paragraph 254, wherein said cell is contacted with said antisense nucleic acid by introducing a promoter adjacent to a chromosomal copy of said antisense nucleic acid such that said promoter directs the synthesis of said antisense nucleic acid.
 - 263. The method of Paragraph 254, wherein said cell is contacted with said antisense nucleic acid by introducing a retron which expresses said antisense nucleic acid into said cell population.
- 264. The method of Paragraph 254, wherein said cell is contacted with said antisense nucleic acid by introducing a ribozyme into said cell-population, wherein a binding portion of said ribozyme is complementary to said antisense oligonucleotide.

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265. The method of Paragraph 254, wherein said cell is contacted with said antisense nucleic acid by introducing a liposome comprising said antisense oligonucleotide into said cell.

- 266. The method of Paragraph 254, wherein said cell is contacted with said antisense nucleic acid by electroporation of said antisense nucleic acid into said cell.
- 267. The method of Paragraph 254, wherein said antisense nucleic acid has at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence comprising at least 10, at least 20, at least 25, at least 30, at least 50 or more than 50 consecutive nucleotides of one of SEO ID NOs.: 8-3795.
- 268. The method of Paragraph 254 wherein said antisense nucleic acid is a synthetic oligonucleotide.
- 269. The method of Paragraph 254, wherein said gene comprises a nucleic acid selected from the group consisting of a nucleic acid comprising a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, a nucleic acid comprising a nucleotide sequence which hybridizes to a sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under moderate conditions.
- 270. A method for identifying a gene which is required for proliferation of a cell comprising:
 - (a) contacting a cell with an antisense nucleic acid selected from the group consisting of a nucleic acid at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 or a proliferation-inhibiting portion thereof, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, wherein said cell is a cell other than the organism from which said nucleic acid was obtained;
 - (b) determining whether said nucleic acid inhibits proliferation of said cell; and
 - (c) identifying the gene in said cell which encodes the mRNA which is complementary to said antisense nucleic acid or a portion thereof.
- 271. The method of Paragraph 270, wherein said cell is selected from the group consisting of *Staphylococcus* species, *Streptococcus* species, *Enterococcus* species, *Mycobacterium* species, *Clostridium* species, and *Bacillus* species.

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272. The method of Paragraph 270 wherein said cell is selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata). Candida tropicalis. Candida parapsilosis. Candida guilliermondii. Candida krusei, Candida kefyr (also called Candida pseudotropicalis). Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corvnebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium. Escherichia coli. Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae. Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori. Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella bovdii. Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

- 273. The method of Paragraph 270, wherein said cell is not E. coli.
- 274. The method of Paragraph 270, further comprising operably linking said antisense
 20 nucleic acid to a promoter which is functional in said cell, said promoter being included in a vector, and introducing said vector into said cell.
 - 275. A method for identifying a compound having the ability to inhibit proliferation of a cell comprising:
 - (a) identifying a homolog of a gene or gene product whose activity or level is inhibited by an antisense nucleic acid in a test cell, wherein said test cell is not the microorgaism from which the antisense nucleic acid was obtained, wherein said antisense nucleic acid is selected from the group consisting of a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOs. 8-3795, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions;
 - (b) identifying an inhibitory nucleic acid sequence which inhibits the activity of said homolog in said test cell;
 - (c) contacting said test cell with a sublethal level of said inhibitory nucleic acid, thus sensitizing said cell;

(d) contacting the sensitized cell of step (c) with a compound; and

- (e) determining the degree to which said compound inhibits proliferation of said sensitized cell relative to a cell which does not express said inhibitory nucleic acid.
- 276. The method of Paragraph 275, wherein said determining step comprises determining whether said compound inhibits proliferation of said sensitized test cell to a greater extent than said compound inhibits proliferation of a nonsensitized test cell.

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- 277. The method of Paragraph 275, wherein step (a) comprises identifying a homologous nucleic acid to a gene or gene product whose activity or level is inhibited by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOs. 8-3795 or a nucleic acid encoding a homologous polypeptide to a polypeptide whose activity or level is inhibited by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOs. 8-3795 by using an algorithm selected from the group consisting of BLASTN version 2.0 with the default parameters and FASTA version 3.0t78 algorithm with the default parameters to identify said homologous nucleic acid or said nucleic acid encoding a homologous polypeptide in a database.
- 278. The method of Paragraph 275 wherein said step (a) comprises identifying a homologous nucleic acid or a nucleic acid encoding a homologous polypeptide by identifying nucleic acids comprising nucleotide sequences which hybridize to said nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOs. 8-3795 or the complement of the nucleotide sequence of said nucleic acid selected from the group consisting of SEQ ID NOs. 8-3795.
- 279. The method of Paragraph 275 wherein step (a) comprises expressing a nucleic acid having at least 70% nucleic acid identity as determined using BLASTN version 2.0 with the default parameters to a sequence selected from the group consisting of SEQ ID NOs. 8-3795 in said test cell.
- 280. The method of Paragraph 275, wherein step (a) comprises identifying a homologous nucleic acid or a nucleic acid encoding a homologous polypeptide in an test cell selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus

jaecaus, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris.

- 5 Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.
 - 281. The method of Paragraph 275, wherein step (a) comprises identifying a homologous nucleic acid or a nucleic acid encoding a homologous polypeptide in a test cell other than *E. coli*.
 - 282. The method of Paragraph 275, wherein said inhibitory nucleic acid is an antisense nucleic acid.

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- 283. The method of Paragraph 275, wherein said inhibitory nucleic acid comprises an antisense nucleic acid to a portion of said homolog.
- 284. The method of Paragraph 275, wherein said inhibitory nucleic acid comprises an antisense nucleic acid to a portion of the operon encoding said homolog.
- 20 285. The method of Paragraph 275, wherein the step of contacting the cell with a sublethal level of said inhibitory nucleic acid comprises directly contacting said cell with said inhibitory nucleic acid.
 - 286. The method of Paragraph 275, wherein the step of contacting the cell with a sublethal level of said inhibitory nucleic acid comprises expressing an antisense nucleic acid to said homolog in said cell.
 - 287. The method of Paragraph 275, wherein said gene product comprises a polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
- from the group consisting of a nucleic acid comprising a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under moderate conditions.

289. A compound identified using the method of Paragraph 275.

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290. A method of identifying a compound having the ability to inhibit proliferation comprising:

- (a) sensitizing a test cell by contacting said test cell with a sublethal level of an antisense nucleic acid, wherein said antisense nucleic acid is selected from the group consisting of a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOs. 8-3795 or a portion thereof which inhibits the proliferation of the cell from which said nucleic acid was obtained, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditionst:
 - (b) contacting the sensitized test cell of step (a) with a compound; and
- (c) determining the degree to which said compound inhibits proliferation of said sensitized test cell relative to a cell which does not contain said antisense nucleic acid.
- 291. The method of Paragraph 290, wherein said determining step comprises determining whether said compound inhibits proliferation of said sensitized test cell to a greater extent than said compound inhibits proliferation of a nonsensitized test cell.
 - 292. A compound identified using the method of Paragraph 290.
- 293. The method of Paragraph 290, wherein said test cell is selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli. Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis. Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica,

Yersinia pestis and any species falling within the genera of any of the above species.

294. The method of Paragraph 290, wherein the test cell is not E. coli.

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295. A method for identifying a compound having activity against a biological pathway required for proliferation comprising:

- (a) sensitizing a cell by providing a sublethal level of an antisense nucleic acid complementary to a nucleic acid encoding a gene product required for proliferation. wherein said gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.; 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEO ID NOs: 8-3795:
 - (b) contacting the sensitized cell with a compound; and
- 25 (c) determining the extent to which said compound inhibits the growth of said sensitized cell relative to a cell which does not contain said antisense nucleic acid.
 - 296. The method of Paragraph 295, wherein said determining step comprises determining whether said compound inhibits the growth of said sensitized cell to a greater extent than said compound inhibits the growth of a nonsensitized cell.
 - 297. The method of Paragraph 295, wherein said cell is selected from the group consisting of bacterial cells, fungal cells, plant cells, and animal cells.
 - 298. The method of Paragraph 295, wherein said cell is a Gram positive bacterium.
 - 299. The method of Paragraph 298, wherein said Gram positive bacterium is selected from the group consisting of *Staphylococcus* species, *Streptococcus* species, *Enterococcus* species, *Mycobacterium* species, *Clostridium* species, and *Bacillus* species.
 - 300. The method of Paragraph 299, wherein said Gram positive bacterium is Staphylococcus aureus.

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301. The method of Paragraph 298, wherein said Gram positive bacterium is selected from the group consisting of *Staphylococcus aureus* RN450 and *Staphylococcus aureus* RN4220.

The method of Paragraph 295, wherein said cell is selected from the group 302. consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae. Enterococcus faecalis, Enterococcus faecium, Escherichia coli. Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae. Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida. Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica. Yersinia pestis and any species falling within the genera of any of the above species.

303. The method of Paragraph 295, wherein said cell is not an E. coli cell.

- 304. The method of Paragraph 295, wherein said gene product is from an organism other than *E. coli*.
- 305. The method of Paragraph 295, wherein said antisense nucleic acid is transcribed from an inducible promoter.
- 306. The method of Paragraph 305, further comprising contacting the cell with an agent which induces expression of said antisense nucleic acid from said inducible promoter, wherein said antisense nucleic acid is expressed at a sublethal level.
- 307. The method of Paragraph 295, wherein inhibition of proliferation is measured by monitoring the optical density of a liquid culture.
- 308. The method of Paragraph 295, wherein said gene product comprises a polypeptide having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
- 309. The method of Paragraph 295, wherein said nucleic acid encoding said gene product comprises a nucleic acid selected from the group consisting of a nucleic acid comprising a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting

of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under moderate condtions.

310. A compound identified using the method of Paragraph 295.

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- 311. A method for identifying a compound having the ability to inhibit cellular proliferation comprising:
 - (a) contacting a cell with an agent which reduces the activity or level of a gene product required for proliferation of said cell, wherein said gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEO ID NOs: 8-3795:
 - (b) contacting said cell with a compound; and
 - (c) determining the degree to which said compound reduces proliferation of said contacted cell relative to a cell which was not contacted with said agent.
- 312. The method of Paragraph 311, wherein said determining step comprises determining whether said compound reduces proliferation of said contacted cell to a greater extent than said compound reduces proliferation of cells which have not been contacted with said agent.
- 313. The method of Paragraph 311, wherein said cell is selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida

giabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

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- 314. The method of Paragraph 311, wherein said cell is not an E. coli cell.
- 315. The method of Paragraph 311, wherein said gene product is from an organism other than *E. coli*.
- 316. The method of Paragraph 311, wherein said agent which reduces the activity or level of a gene product required for proliferation of said cell comprises an antisense nucleic acid to a gene or operon required for proliferation.
- 317. The method of Paragraph 311, wherein said agent which reduces the activity or level of a gene product required for proliferation of said cell comprises a compound known to inhibit growth or proliferation of a cell.
- 318. The method of Paragraph 311, wherein said cell contains a mutation which reduces the activity or level of said gene product required for proliferation of said cell.
- 319. The method of Paragraph 311, wherein said mutation is a temperature sensitive mutation.
- 320. The method of Paragraph 311, wherein said gene product comprises a gene product comprises a polypeptide having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
 - 321. A compound identified using the method of Paragraph 311.
- 322. A method for identifying the biological pathway in which a proliferation-required gene product or a gene encoding a proliferation-required gene product lies comprising:
 - (a) providing a sublethal level of an antisense nucleic acid which inhibits the activity or reduces the level of said gene encoding a proliferation-required gene product or said said proliferation-required gene product in a test cell, wherein said proliferation-

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required gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEO ID NOs: 8-3795:

(b) contacting said test cell with a compound known to inhibit growth or proliferation of a cell, wherein the biological pathway on which said compound acts is known; and

- (c) determining the degree to which said compound inhibits proliferation of said test cell relative to a cell which does not contain said antisense nucleic acid.
- 323. The method of Paragraph 322, wherein said determining step comprises determining whether said test cell has a substantially greater sensitivity to said compound than a cell which does not express said sublethal level of said antisense nucleic acid.
- 324. The method of Paragraph 322, wherein said gene product comprises a polypeptide having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
- 325. The method of Paragraph 322, wherein said test cell is selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus

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neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species.

- 326. The method of Paragraph 322, wherein said test cell is not an E. coli cell.
- 327. The method of Paragraph 322, wherein said gene product is from an organism other than E. coli.
- 328. A method for determining the biological pathway on which a test compound acts comprising:
 - (a) providing a sublethal level of an antisense nucleic acid complementary to a proliferation-required nucleic acid in a cell, thereby producing a sensitized cell, wherein said antisense nucleic acid is selected from the group consisting of a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795 or a proliferation-inhibiting portion thereof, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions and wherein the biological pathway in which said proliferation-required nucleic acid or a protein encoded by said proliferation-required polypeptide lies is known,
 - (b) contacting said cell with said test compound; and
 - (c) determining the degree to which said compound inhibits proliferation of said sensitized cell relative to a cell which does not contain said antisense nucleic acid.
- 329. The method of Paragraph 328, wherein said determining step comprises determining whether said sensitized cell has a substantially greater sensitivity to said test compound than a cell which does not express said sublethal level of said antisense nucleic acid.
 - 330. The method of Paragraph 328, further comprising:
- (d) providing a sublethal level of a second antisense nucleic acid complementary to a second proliferation-required nucleic acid in a second cell, wherein said second

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proliferation-required nucleic acid is in a different biological pathway than said proliferation-required nucleic acid in step (a); and

- (e) determining whether said second cell does not have a substantially greater sensitivity to said test compound than a cell which does not express said sublethal level of said second antisense nucleic acid, wherein said test compound is specific for the biological pathway against which the antisense nucleic acid of step (a) acts if said sensitized cell has substantially greater sensitivity to said test compound than said second cell.
- 331. The method of Paragraph 328, wherein said sensitized cell is selected from the group consisting of Anaplasma marginale, Aspergillus fimigatus, Bacillus anthracis, Bacterioides 10 fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans. Candida glabrata (also called Torulopsis glabrata), Candida tropicalis. Candida parapsilosis. Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, 15 Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, 20 Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of 25 the above species.
 - 332. The method of Paragraph 328, wherein said sensitized cell is not an E. coli cell.
 - 333. The method of Paragraph 328, wherein said proliferation-required nucleic acid is from an organism other than *E. coli*.
 - 334. A compound which inhibits proliferation by interacting with a gene encoding a gene product required for proliferation or with a gene product required for proliferation, wherein said gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from

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the group consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795.

- 335. The compound of Paragraph 334, wherein said gene product comprises a polypeptide having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
- 336. The compound of Paragraph 334, wherein said gene comprises a nucleic acid selected from the group consisting of a nucleic acid comprising a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 under moderate conditions.
 - 337. A method for manufacturing an antibiotic comprising the steps of: screening one or more candidate compounds to identify a compound that reduces the

screening one or more candidate compounds to identify a compound that reduces the activity or level of a gene product required for proliferation wherein said gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence

which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795; and

manufacturing the compound so identified.

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- 338. The method of Paragraph 337, wherein said screening step comprises performing any one of the methods of Paragraphs 205, 211, 222, 275, 290, 295, 311.
- 339. The method of Paragraph 337, wherein said gene product comprises a polypeptide having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.
- 340. A method for inhibiting proliferation of a cell in a subject comprising administering an effective amount of a compound that reduces the activity or level of a gene product required for proliferation of said cell, wherein said gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795.
- 341. The method of Paragraph 340 wherein said subject is selected from the group consisting of vertebrates, mammals, avians, and human beings.
- 342. The method of Paragraph 340, wherein said gene product comprises a polypeptide having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default

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parameters to an amino acid sequence selected from the group consisting of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110.

- 343. The method of Paragraph 340, wherein said cell is selected from the group consisting of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata). Candida tropicalis, Candida parapsilosis. Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica. Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori. Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica. Yersinia pestis and any species falling within the genera of any of the above species.
 - 344. The method of Paragraph 340, wherein said cell is not E. coli.
- 345. The method of Paragraph 340, wherein said gene product is from an organism other than *E. coli*.

Definitions

By "biological pathway" is meant any discrete cell function or process that is carried out by a gene product or a subset of gene products. Biological pathways include anabolic, catabolic, enzymatic, biochemical and metabolic pathways as well as pathways involved in the production of cellular structures such as cell walls. Biological pathways that are usually required for proliferation of cells or microorganisms include, but are not limited to, cell division, DNA synthesis and replication, RNA synthesis (transcription), protein synthesis (translation), protein processing, protein transport, fatty acid biosynthesis, electron transport chains, cell wall synthesis, cell membrane production, synthesis and maintenance, and the like.

By "inhibit activity of a gene or gene product" is meant having the ability to interfere with the function of a gene or gene product in such a way as to decrease expression of the gene, in such a way as to reduce the level or activity of a product of the gene or in such a way as to inhibit the interaction of the gene or gene product with other biological molecules required for its activity. Agents which inhibit the activity of a gene include agents that inhibit transcription of the gene, agents that inhibit processing of the transcript of the gene, agents that reduce the stability of the

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transcript of the gene, and agents that inhibit translation of the mRNA transcribed from the gene. In microorganisms, agents which inhibit the activity of a gene can act to decrease expression of the operon in which the gene resides or alter the folding or processing of operon RNA so as to reduce the level or activity of the gene product. The gene product can be a non-translated RNA such as ribosomal RNA, a translated RNA (mRNA) or the protein product resulting from translation of the gene mRNA. Of particular utility to the present invention are antisense RNAs that have activities against the operons or genes to which they specifically hybridze.

By "activity against a gene product" is meant having the ability to inhibit the function or to reduce the level or activity of the gene product in a cell. This includes, but is not limited to, inhibiting the enzymatic activity of the gene product or the ability of the gene product to interact with other biological molecules required for its activity, including inhibiting the gene product's assembly into a multimeric structure.

By "activity against a protein" is meant having the ability to inhibit the function or to reduce the level or activity of the protein in a cell. This includes, but is not limited to, inhibiting the enzymatic activity of the protein or the ability of the protein to interact with other biological molecules required for its activity, including inhibiting the protein's assembly into a multimeric structure.

By "activity against a nucleic acid" is meant having the ability to inhibit the function or to reduce the level or activity of the nucleic acid in a cell. This includes, but is not limited to, inhibiting the ability of the nucleic acid interact with other biological molecules required for its activity, including inhibiting the nucleic acid's assembly into a multimeric structure.

By "activity against a gene" is meant having the ability to inhibit the function or expression of the gene in a cell. This includes, but is not limited to, inhibiting the ability of the gene to interact with other biological molecules required for its activity.

By "activity against an operon" is meant having the ability to inhibit the function or reduce the level of one or more products of the operon in a cell. This includes, but is not limited to, inhibiting the enzymatic activity of one or more products of the operon or the ability of one or more products of the operon to interact with other biological molecules required for its activity.

By "antibiotic" is meant an agent which inhibits the proliferation of a cell or microorganism.

By "E. coli or Escherichia coli" is meant Escherichia coli or any organism previously categorized as a species of Shigella including Shigella boydii, Shigella flexneri, Shigella dysenteriae, Shigella sonnei, Shigella 2A.

By "homologous coding nucleic acid" is meant a nucleic acid homologous to a nucleic acid encoding a gene product whose activity or level is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 or a portion thereof. In some embodiments, the homologous coding nucleic acid may have at least 97%, at least 95%, at least 90%, at least 85%, at

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least 80%, or at least 70% nucleotide sequence identity to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 and fragments comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides thereof. In other embodiments the homologous coding nucleic acids may have at least 97%, at least 95%, at least 90%, at least 85%, at least 80%, or at least 70% nucleotide sequence identity to a nucleotide sequence selected from the group consisting of the nucleotide sequences complementary to one of SEQ ID NOs.: 8-3795 and fragments comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75. 100, 150, 200, 300, 400, or 500 consecutive nucleotides thereof. Identity may be measured using BLASTN version 2.0 with the default parameters or tBLASTX with the default parameters. (Altschul, S.F. et al. Gapped BLAST and PSI-BLAST: A New Generation of Protein Database Search Programs, Nucleic Acid Res. 25: 3389-3402 (1997)) Alternatively a "homologuous coding nucleic acid" could be identified by membership of the gene of interest to a functional orthologue cluster. All other members of that orthologue cluster would be considered homologues. Such a library of functional orthologue clusters can be found at http://www.ncbi.nlm.nih.gov/COG. A gene can be classified into a cluster of orthologous groups or COG by using the COGNITOR program available at the above web site, or by direct BLASTP comparison of the gene of interest to the members of the COGs and analysis of these results as described by Tatusov, R.L., Galperin. M.Y., Natale, D. A. and Koonin, E.V. (2000) The COG database; a tool for genome-scale analysis of protein functions and evolution. Nucleic Acids Research v. 28 n. 1, pp33-36.

The term "homologous coding nucleic acid" also includes nucleic acids comprising nucleotide sequences which encode polypeptides having at least 99%, 95%, at least 90%, at least 85%, at least 80%, at least 70%, at least 60%, at least 50%, at least 40% or at least 25% maino acid identity or similarity to a polypeptide comprising the amino acid sequence of one of SEQ ID NOs: 3801-3805, 4861-5915, 10013-14110 or to a polypeptide whose expression is inhibited by a nucleic acid comprising a nucleotide sequence of one of SEQ ID NOs: 8-3795 or fragments comprising at least 5, 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, or 150 consecutive amino acids thereof as determined using the FASTA version 3.0t78 algorithm with the default parameters.

Alternatively, protein identity or similarity may be identified using BLASTP with the default parameters, or tBLASTX with the default parameters, TBLASTN with the default parameters, or tBLASTX with the default parameters. (Altschul, S.F. et al. Gapped BLAST and PSI-BLAST: A New Generation of Protein Database Search Programs, Nucleic Acid Res. 25: 3389-3402 (1997)).

The term "homologous coding nucleic acid" also includes coding nucleic acids which hybridize under stringent conditions to a nucleic acid selected from the group consisting of the nucleotide sequences complementary to one of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 and coding nucleic acids comprising nucleotide sequences which hybridize under stringent conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of the sequences complementary to one of SEQ ID NOS.:

3796-3800, 3806-4860, 5916-10012 As used herein, "stringent conditions" means hybridization to filter-bound nucleic acid in 6xSSC at about 45°C followed by one or more washes in 0.1xSSC/0.2% SDS at about 68°C. Other exemplary stringent conditions may refer, *e.g.*, to washing in 6xSSC/0.05% sodium pyrophosphate at 37°C, 48°C, 55°C, and 60°C as appropriate for the particular probe being used.

The term "homologous coding nucleic acid" also includes coding nucleic acids comprising nucleotide sequences which hybridize under moderate conditions to a nucleotide sequence selected from the group consisting of the sequences complementary to one of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 and coding nucleic acids comprising nucleotide sequences which hybridize under moderate conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of the sequences complementary to one of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012. As used herein, "moderate conditions" means hybridization to filter-bound DNA in 6x sodium chloride/sodium citrate (SSC) at about 45°C followed by one or more washes in 0.2xSSC/0.1% SDS at about 42-65°C.

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The term "homologous coding nucleic acids" also includes nucleic acids comprising nucleotide sequences which encode a gene product whose activity may be complemented by a gene encoding a gene product whose activity is inhibited by a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795. In some embodiments, the homologous coding nucleic acids may encode a gene product whose activity is complemented by the gene product encoded by a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012. In other embodiments, the homologous coding nucleic acids may comprise a nucleotide sequence encode a gene product whose activity is complemented by one of the polypeptides of SEQ ID NOs. 3745-4773.

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The term "homologous antisense nucleic acid" includes nucleic acids comprising a nucleotide sequence having at least 97%, at least 95%, at least 90%, at least 85%, at least 80%, or at least 70% nucleotide sequence identity to a nucleotide sequence selected from the group consisting of one of the sequences of SEQ ID NOS. 8-3795 and fragments comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides thereof. Homologous antisense nucleic acids may also comprising nucleotide sequences which have at least 97%, at least 95%, at least 90%, at least 85%, at least 80%, or at least 70% nucleotide sequence identity to a nucleotide sequence selected from the group consisting of the sequences complementary to one of sequences of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 and fragments comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides thereof. Nucleic acid identity may be determined as described above.

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The term "homologous antisense nucleic acid" also includes antisense nucleic acids comprising nucleotide sequences which hybridize under stringent conditions to a nucleotide sequence complementary to one of SEQ ID NOs.: 8-3795 and antisens nucleic acids comprising

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nucleotide sequences which hybridize under stringent conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of the sequence complementary to one of SEQ ID NOs. 8-3795. Homologous antisense nucleic acids also include antisense nucleic acids comprising nucleotide sequences which hybridize under stringent conditions to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 and antisense nucleic acids comprising nucleotide sequences which hybridize under stringent conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of one of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012.

The term "homologous antisense nucleic acid" also includes antisense nucleic acids comprising nucleotide sequences which hybridize under moderate conditions to a nucleotide sequence complementary to one of SEQ ID NOs.: 8-3795 and antisens nucleic acids comprising nucleotide seuqences which hybridize under moderate conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of the sequence complementary to one of SEQ ID NOs. 8-3795. Homologous antisense nucleic acids also include antisense nucleic acids comprising nucleotide seuqences which hybridize under moderate conditions to a nucleotide sequence selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 and antisense nucleic acids which comprising nucleotide sequences hybridize under moderate conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of one of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012.

By "homologous polypeptide" is meant a polypeptide homologous to a polypeptide whose activity or level is inhibited by a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 or by a homologous antisense nucleic acid. The term "homologous polypeptide" includes polypeptides having at least 99%, 95%, at least 90%, at least 85%, at least 80%, at least 70%, at least 60%, at least 50%, at least 40% or at least 25% amino acid identity or similarity to a polypeptide whose activity or level is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795 or by a homologous antisense nucleic acid, or polypeptides having at least 99%, 95%, at least 90%, at least 85%, at least 80%, at least 70%, at least 60%, at least 50%, at least 40% or at least 25% amino acid identity or similarity to a polypeptide to a fragment comprising at least 5, 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, or 150 consecutive amino acids of a polypeptide whose activity or level is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 or by a homologous antisense nucleic acid. Identity or similarity may be determined using the FASTA version 3.0t78 algorithm with the default parameters. Alternatively, protein identity or similarity may be identified using BLASTP with the default parameters, or TBLASTN with the default

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parameters. (Altschul, S.F. et al. Gapped BLAST and PSI-BLAST: A New Generation of Protein Database Search Programs, Nucleic Acid Res. 25: 3389-3402 (1997).

The term homologous polypeptide also includes polypeptides having at least 99%, 95%, at least 90%, at least 85%, at least 80%, at least 70%, at least 60%, at least 50%, at least 40% or at least 25% amino acid identity or similarity to a polypeptide selected from the group consisting of SEQ ID NOs: 3801-3805, 4861-5915, 10013-14110 and polypeptides having at least 99%, 95%, at least 90%, at least 85%, at least 80%, at least 70%, at least 60%, at least 50%, at least 40% or at least 25% amino acid identity or similarity to a fragment comprising at least 5, 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, or 150 consecutive amino acids of a polypeptide selected from the group consisting of SEQ ID NOs: 3801-3805, 4861-5915, 10013-14110.

The invention also includes polynucleotides, preferably DNA molecules, that hybridize to one of the nucleic acids of SEQ ID NOs.: 8-3795, SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012 or the complements of any of the preceding nucleic acids. Such hybridization may be under stringent or moderate conditions as defined above or under other conditions which permit specific hybridization. The nucleic acid molecules of the invention that hybridize to these DNA sequences include oligodeoxynucleotides ("oligos") which hybridize to the target gene under highly stringent or stringent conditions. In general, for oligos between 14 and 70 nucleotides in length the melting temperature (Tm) is calculated using the formula:

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$$\text{Tm}(^{\circ}\text{C}) = 81.5 + 16.6(\log[\text{monovalent cations (molar)}] + 0.41 (\% G+C) - (500/N)$$

where N is the length of the probe. If the hybridization is carried out in a solution containing formamide, the melting temperature may be calculated using the equation:

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$$Tm(^{\circ}C) = 81.5 + 16.6(log[monovalent cations (molar)] + 0.41(% G+C) - (0.61)$$

(% formamide) - (500/N)

where N is the length of the probe. In general, hybridization is carried out at about 20-25 degrees below Tm (for DNA-DNA hybrids) or about 10-15 degrees below Tm (for RNA-DNA hybrids).

Other hybridization conditions are apparent to those of skill in the art (see, for example, Ausubel, F.M. et al., eds., 1989, Current Protocols in Molecular Biology, Vol. I, Green Publishing Associates, Inc. and John Wiley & Sons, Inc., New York, at pp. 6.3.1-6.3.6 and 2.10.3.

The term, *Salmonella*, is the generic name for a large group of gram-negative enteric bacteria that are closely related to *Escherichia coli*. The diseases caused by *Salmonella* are often due to contamination of foodstuffs or the water supply and affect millions of people each year. Traditional methods of *Salmonella* taxonomy were based on assigning a separate species name to

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each serologically distinguishable strain (Kauffmann, F 1966 The bacteriology of the *Enterobacteriaceae*. Munksgaard, Copenhagen). Serology of *Salmonella* is based on surface antigens (O [somatic] and H [flagellar]). Over 2,400 serotypes or serovars of *Salmonella* are known (Popoff, et al. 2000 Res. Microbiol. 151:63-65). Therefore, each serotype was considered to be a separate species and often given names, accordingly (e.g. *S. paratyphi, S. typhimurium, S. typhi. S. enteriditis*, etc.).

However, by the 1970s and 1980s it was recognized that this system was not only cumbersome, but also inaccurate. Then, many *Salmonella* species were lumped into a single species (all serotypes and subgenera I, II, and IV and all serotypes of *Arizona*) with a second subspecies, *S. bongorii* also recognized (Crosa, et al., 1973, J. Bacteriol. 115:307-315). Though species designations are based on the highly variable surface antigens, the *Salmonella* are very similar otherwise with a major exception being pathogenicity determinants.

There has been some debate on the correct name for the Salmonella species. Currently (Brenner, et al. 2000 J. Clin. Microbiol. 38:2465-2467), the accepted name is Salmonella enterica. S. enterica is divided into six subspecies (I, S. enterica subsp. enterica; II, S. enterica, subsp. salamae; IIIa, S. enterica subsp. arizonàe; IIIb, S. enterica subsp. diarizonae; IV, S. enterica subsp. houtenae; and VI, S. enterica subsp. indica). Within subspecies I, serotypes are used to distinguish each of the serotypes or serovars (e.g. S. enterica serotype Enteriditis, S. enterica serotype Typhimurium, S. enterica serotype Typhi, and S. enterica serotype Choleraesuis, etc.). Current convention is to spell this out on first usage (Salmonella enterica ser. Typhimurium) and then use an abbreviated form (Salmonella Typhimurium or S. Typhimurium). Note, the genus and species names (Salmonella enterica) are italicized but not the serotype/serovar name (Typhimurium). Because the taxonomic committees have yet to officially approve of the actual species name, this latter system is what is employed by the CDC (Brenner, et al. 2000 J. Clin. Microbiol. 38:2465-2467). Due to the concerns of both taxonomic priority and medical importance, some of these serotypes might ultimately receive full species designations (S.typhi would be the most notable).

Therefore, as used herein "Salmonella enterica" includes serovars Typhi, Typhimurium, Paratyphi, Choleraesuis, etc." However, appeals of the "official" name are in process and the taxonomic designations may change (S. choleraesuis is the species name that could replace S. enterica based solely on priority).

By "identifying a compound" is meant to screen one or more compounds in a collection of compounds such as a combinatorial chemical library or other library of chemical compounds or to characterize a single compound by testing the compound in a given assay and determining whether it exhibits the desired activity.

By "inducer" is meant an agent or solution which, when placed in contact with a cell or microorganism, increases transcription, or inhibitor and/or promoter clearance/fidelity, from a desired promoter.

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As used herein, "nucleic acid" means DNA, RNA, or modified nucleic acids. Thus, the terminology "the nucleic acid of SEQ ID NO: X" or "the nucleic acid comprising the nucleotide sequence" includes both the DNA sequence of SEQ ID NO: X and an RNA sequence in which the thymidines in the DNA sequence have been substituted with uridines in the RNA sequence and in which the deoxyribose backbone of the DNA sequence has been substituted with a ribose backbone in the RNA sequence. Modified nucleic acids are nucleic acids having nucleotides or structures which do not occur in nature, such as nucleic acids in which the internucleotide phosphate residues with methylphosphonates, phosphorothioates, phosphoramidates, and phosphate esters.

Nonphosphate internucleotide analogs such as siloxane bridges, carbonate brides, thioester bridges, as well as many others known in the art may also be used in modified nucleic acids. Modified nucleic acids may also comprise, α -anomeric nucleotide units and modified nucleotides such as 1,2-dideoxy-d-ribofuranose, 1,2-dideoxy-1-phenylribofuranose, and N^4 , N^4 -ethano-5-methyl-cytosine are contemplated for use in the present invention. Modified nucleic acids may also be peptide nucleic acids in which the entire deoxyribose-phosphate backbone has been exchanged with a chemically completely different, but structurally homologous, polyamide (peptide) backbone containing 2-aminoethyl glycine units.

As used herein, "sub-lethal" means a concentration of an agent below the concentration required to inhibit all cell growth.

Brief Description of the Drawings

Figure 1 is an IPTG dose response curve in *E. coli* transformed with an IPTG-inducible plasmid containing either an antisense clone to the *E. coli* ribosomal protein *rplW* (AS-*rplW*) which is required for protein synthesis and essential for cell proliferation, or an antisense clone to the *elaD* (AS-*elaD*) gene which is not known to be involved in protein synthesis and which is also essential for proliferation.

Figure 2A is a tetracycline dose response curve in *E. coli* transformed with an IPTG-inducible plasmid containing antisense to *rplW* (AS-*rplW*) in the absence (0) or presence of IPTG at concentrations that result in 20% and 50% growth inhibition.

Figure 2B is a tetracycline dose response curve in *E. coli* transformed with an IPTG-inducible plasmid containing antisense to *elaD* (AS-*elaD*)in the absence (0) or presence of IPTG at concentrations that result in 20% and 50% growth inhibition.

Figure 3 is a graph showing the fold increase in tetracycline sensitivity of *E. coli* transfected with antisense clones to essential ribosomal proteins *L23* (AS-*rplW*) and *L7/L12* and *L10* (AS-*rplLrplJ*). Antisense clones to genes known to not be directly involved in protein synthesis, *atpB/E* (AS-*atpB/E*), *visC* (AS-*visC*), *elaD* (AS-*elaD*), *yohH* (AS-*yohH*), are much less sensitive to tetracycline.

Figure 4 illustrates the results of an assay in which *Staphylococcus aureus* cells transcribing an antisense nucleic acid complementary to the *gyrB* gene encoding the β subunit of gyrase were contacted with several antibiotics whose targets were known.

Detailed Description of the Preferred Embodiments

The present invention describes a group of prokaryotic genes and gene families required for cellular proliferation. Exemplary genes and gene families from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae. Pseudomonas aeruginosa. Staphylococcus aureus. and Salmonella typhi are provided. A proliferation-required gene or gene family is one where, in the absence or substantial reduction of a gene transcript and/or gene product, growth or viability of the cell or microorganism is reduced or eliminated. Thus, as used herein, the terminology "proliferation-required" or "required for proliferation" encompasses instances where the absence or substantial reduction of a gene transcript and/or gene product completely eliminates cell growth as well as instances where the absence of a gene transcript and/or gene product merely reduces cell growth. These proliferation-required genes can be used as potential targets for the generation of new antimicrobial agents. To achieve that goal, the present invention also encompasses assays for analyzing proliferation-required genes and for identifying compounds which interact with the gene and/or gene products of the proliferation-required genes. In addition, the present invention contemplates the expression of genes and the purification of the proteins encoded by the nucleic acid sequences identified as required proliferation genes and reported herein. The purified proteins can be used to generate reagents and screen small molecule libraries or other candidate compound libraries for compounds that can be further developed to yield novel antimicrobial compounds.

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The present invention also describes methods for identification of nucleotide sequences homologous to these genes and polypeptides described herein, including nucleic acids comprising nucleotide sequences homologous to the nucleic acids of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 and polypeptides homologous to the polypeptides of SEQ ID NOS.: 3801-3805, 4861-5915, 10013-14110. For example, these sequences may be used to identify homologous coding nucleic acids, homologous antisense nucleic acids, or homologous polypeptides in microorganisms such as Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli,

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Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis or any species falling within the genera of any of the above species. In some embodiments, the homologous coding nucleic acids, homologus antisense nucleic acids, or homologous polypeptides are identified in an organism other than E. coli.

The homologous coding nucleic acids, homologous antisense nucleic acids, or homologous polypeptides, may then be used in each of the methods described herein, including methods to identify compounds which inhibit the proliferation of the organism containing the homologous coding nucleic acid, homologous antisense nucleic acid or homologous polypeptide, methods of inhibiting the growth of the organism containing the homologous coding nucleic acid, homologus antisense nucleic acid or homologous polypeptide, methods of identifying compounds which influence the activity or level of a gene product required for proliferation of the organism containing the homologous coding nucleic acid, homologous antisense nucleic acid or homologous polypeptide, methods for identifying compounds or nucleic acids having the ability to reduce the level or activity of a gene product required for proliferation of the organism containing the homologous coding nucleic acid, homologous antisense nucleic acid or homologous polypeptide, methods of inhibiting the activity or expression of a gene in an operon required for proliferation of the organism containing the homologous coding nucleic acid, homologous antisense nucleic acid or homologous polypeptide, methods for identifying a gene required proliferation of the organism containing the homologous coding nucleic acid, homologous antisense nucleic acid or homologous polypeptide, methods for identifying the biological pathway in which a gene or gene product required for proliferation of the organism containing the homologous coding nucleic acid, homologous antisense nucleic acid or homologous polypeptide lies, methods for identifying compounds having activity against biological pathway required for proliferation of the organism containing the homologous coding nucleic acid, homologous antisense nucleic acid or homologous polypeptide, methods for determining the biological pathway on which a test compound acts, and methods of inhibiting the proliferation of the organism containing the homologous coding nucleic acid, homologous antisense nucleic acid or homologous polypeptide in a subject. In some embodiments of the present invention, the methods are performed using an organism, other than E. coli or a gene or gene product from an organism other than E. coli.

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The present invention utilizes a novel method to identify proliferation-required sequences. Generally, a library of nucleic acid sequences from a given source are subcloned or otherwise inserted immediately downstream of an inducible promoter on an appropriate vector, such as a Staphylococcus aureus/E. coli or Pseudomonas aeruginosa/E. coli shuttle vector, or a vector which will replicate in both Salmonella typhimurium and Klebsiella pneumoniae, or other vector or shuttle vector capable of functioning in the intended organism., thus forming an expression library. It is generally preferred that expression is directed by a regulatable promoter sequence such that expression level can be adjusted by addition of variable concentrations of an inducer molecule or of an inhibitor molecule to the medium. Temperature activated promoters, such as promoters regulated by temperature sensitive repressors. such as the lambda C₁₈₅₇ repressor, are also envisioned. Although the insert nucleic acids may be derived from the chromosome of the cell or microorganism into which the expression vector is to be introduced, because the insert is not in its natural chromosomal location, the insert nucleic acid is an exogenous nucleic acid for the purposes of the discussion herein. The term "expression" is defined as the production of a sense or antisense RNA molecule from a gene, gene fragment, genomic fragment, chromosome, operon or portion thereof. Expression can also be used to refer to the process of peptide or polypeptide synthesis. An expression vector is defined as a vehicle by which a ribonucleic acid (RNA) sequence is transcribed from a nucleic acid sequence carried within the expression vehicle. The expression vector can also contain features that permit translation of a protein product from the transcribed RNA message expressed from the exogenous nucleic acid sequence carried by the expression vector. Accordingly, an expression vector can produce an RNA molecule as its sole product or the expression vector can produce a RNA molecule that is ultimately translated into a protein product.

Once generated, the expression library containing the exogenous nucleic acid sequences is introduced into a population of cells (such as the organism from which the exogenous nucleic acid sequences were obtained) to search for genes that are required for bacterial proliferation. Because the library molecules are foreign, in context, to the population of cells, the expression vectors and the nucleic acid segments contained therein are considered exogenous nucleic acid.

Expression of the exogenous nucleic acid fragments in the test population of cells containing the expression library is then activated. Activation of the expression vectors consists of subjecting the cells containing the vectors to conditions that result in the expression of the exogenous nucleic acid sequences carried by the expression library. The test population of cells is then assayed to determine the effect of expressing the exogenous nucleic acid fragments on the test population of cells. Those expression vectors that negatively impacted the growth of the cells upon induction of expression of the random sequences contained therein were identified, isolated, and purified for further study.

A variety of assays are contemplated to identify nucleic acid sequences that negatively impact growth upon expression. In one embodiment, growth in cultures expressing exogenous nucleic acid sequences and growth in cultures not expressing these sequences is compared. Growth measurements

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are assayed by examining the extent of growth by measuring optical densities. Alternatively, enzymatic assays can be used to measure bacterial growth rates to identify exogenous nucleic acid sequences of interest. Colony size, colony morphology, and cell morphology are additional factors used to evaluate growth of the host cells. Those cultures that fail to grow or grow at a reduced rate under expression conditions are identified as containing an expression vector encoding a nucleic acid fragment that negatively affects a proliferation-required gene.

Once exogenous nucleic acids of interest are identified, they are analyzed. The first step of the analysis is to acquire the nucleotide sequence of the nucleic acid fragment of interest. To achieve this end, the insert in those expression vectors identified as containing a nucleotide sequence of interest is sequenced, using standard techniques well known in the art. The next step of the process is to determine the source of the nucleotide sequence. As used herein "source" means the genomic region containing the cloned fragment.

Determination of the gene(s) corresponding to the nucleotide sequence was achieved by comparing the obtained sequence data with databases containing known protein and nucleotide sequences from various microorganisms. Thus, initial gene identification was made on the basis of significant sequence similarity or identity to either characterized or predicted Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa or Enterococcus faecalis genes or their encoded proteins and/or homologues in other species.

The number of nucleotide and protein sequences available in database systems has been 20 growing exponentially for years. For example, the complete nucleotide sequences of Caenorhabditis elegans and several bacterial genomes, including E. coli, Aeropyrum pernix, Aquifex aeolicus, Archaeoglobus fulgidus, Bacillus subtilis, Borrelia burgdorferi, Chlamydia pneumoniae, Chlamydia trachomatis. Clostridium tetani. Corynebacterium diptheria. Deinococcus radiodurans. Haemophilus influenzae, Helicobacter pylori 26695, Helicobacter pylori J99, Methanobacterium 25 thermoautotrophicum, Methanococcus jannaschii, Mycobacterium tuberculosis, Mycoplasma genitalium, Mycoplasma pneumoniae, Pseudomonas aeruginosa, Pyrococcus abyssi, Pyrococcus horikoshii, Rickettsia prowazekii, Synechocystis PCC6803, Thermotoga maritima, Treponema pallidum, Bordetella pertussis, Campylobacter jejuni, Clostridium acetobutylicum, Mycobacterium tuberculosis CSU#93, Neisseria gonorrhoeae, Neisseria meningitidis, Pseudomonas aeruginosa, 30 Pyrobaculum aerophilum, Pyrococcus furiosus, Rhodobacter capsulatus, Salmonella typhimurium. Streptococcus mutans, Streptococcus pyogenes, Ureaplasma urealyticum and Vibrio cholera are available. This nucleotide sequence information is stored in a number of databanks, such as GenBank. the National Center for Biotechnology Information (NCBI), the Genome Sequencing Center (http://genome.wustl.edu/gsc/salmonella.shtml).and the Sanger Centre 35 (http://www.sanger.ac.uk/projects/S typhi)which are publicly available for searching. A variety

of computer programs are available to assist in the analysis of the sequences stored within these databases. FASTA, (W. R. Pearson (1990) "Rapid and Sensitive Sequence Comparison with

FASTP and FASTA" Methods in Enzymology 183:63-98), Sequence Retrieval System (SRS), (Etzold & Argos, SRS an indexing and retrieval tool for flat file data libraries. Comput. Appl. Biosci. 9:49-57, 1993) are two examples of computer programs that can be used to analyze sequences of interest. In one embodiment of the present invention, the BLAST family of computer programs, which includes BLASTN version 2.0 with the default parameters, or BLASTX version 2.0 with the default parameters, is used to analyze nucleotide sequences.

BLAST, an acronym for "Basic Local Alignment Search Tool," is a family of programs for database similarity searching. The BLAST family of programs includes: BLASTN, a nucleotide sequence database searching program, BLASTX, a protein database searching program where the input is a nucleic acid sequence; and BLASTP, a protein database searching program. BLAST programs embody a fast algorithm for sequence matching, rigorous statistical methods for judging the significance of matches, and various options for tailoring the program for special situations. Assistance in using the program can be obtained by e-mail at blast@ncbi.nlm.nih.gov. tBLASTX can be used to translate a nucleotide sequence in all three potential reading frames into an amino acid sequence.

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Bacterial genes are often transcribed in polycistronic groups. These groups comprise operons, which are a collection of genes and intergenic sequences under common regulation. The genes of an operon are transcribed on the same mRNA and are often related functionally. Given the nature of the screening protocol, it is possible that the identified exogenous nucleic acid corresponds to a gene or portion thereof with or without adjacent noncoding sequences, an intragenic sequence (i.e. a sequence within a gene), an intergenic sequence (i.e. a sequence between genes), a nucleotide sequence spanning at least a portion of two or more genes, a 5' noncoding region or a 3' noncoding region located upstream or downstream from the actual nucleotide sequence that is required for bacterial proliferation. Accordingly, it is often desirable to determine which gene(s) that is encoded within the operon is individually required for proliferation.

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In one embodiment of the present invention, an operon is identified and then dissected to determine which gene or genes are required for proliferation. Operons can be identified by a variety of means known to those in the art. For example, the RegulonDB DataBase described by Huerta et al. (Nucl. Acids Res. 26:55-59, 1998), which may also be found on the website http://www.cifn.unam.mx/Computational_Biology/regulondb/, provides information about operons in Escherichia coli. The Subtilist database (http://bioweb.pasteur.fr/GenoList/SubtiList), (Moszer, I., Glaser, P. and Danchin, A. (1995) Microbiology 141: 261-268 and Moszer, I (1998) FEBS Letters 430: 28-36), may also be used to predict operons. This database lists genes from the fully sequenced, Gram-positive bacteria, Bacillus subtilis, together with predicted promoters and terminator sites. This information can be used in conjunction with the Staphylococcus aureus genomic sequence data to predict operons and thus produce a list of the genes affected by the antisense nucleic acids of the present invention. The Pseudomonas aeruginosa web site (http://www.pseudomonas.com) can be used to help predict operon organization in this bacterium.

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The databases available from the Genome Sequencing Center (http://genome.wustl.edu/gsc/salmonella.shtml), and the Sanger Centre (http://www.sanger.ac.uk/projects/S___typhi) may be used to predict operons in Salmonella typhimurium. The TIGR microbial database has an incomplete version of the E. faecalis genome http://www.tigr.org/cgi-bin/BlastSearch/blast.cgi?organism=e_faecalis. One can take a nucleotide sequence and BLAST it for homologs.

A number of techniques that are well known in the art can be used to dissect the operon.

Analysis of RNA transcripts by Northern blot or primer extension techniques are commonly used to analyze operon transcripts. In one aspect of this embodiment, gene disruption by homologous recombination is used to individually inactivate the genes of an operon that is thought to contain a gene required for proliferation.

Several gene disruption techniques have been described for the replacement of a functional gene with a mutated, non-functional (null) allele. These techniques generally involve the use of homologous recombination. One technique using homologous recombination in *Staphylococcus aureus* is described in Xia et a.. 1999, Plasmid 42: 144-149. This technique uses crossover PCR to create a null allele with an in-frame deletion of the coding region of a target gene. The null allele is constructed in such a way that nucleotide sequences adjacent to the wild type gene are retained. These homologous sequences surrounding the deletion null allele provide targets for homologous recombination so that the wild type gene on the *Staphylococcus aureus* chromosome can be replaced by the constructed null allele. This method can be used with other bacteria as well, including *Salmonella* and *Klebsiella* species. Similar gene disruption methods that employ the counter selectable marker *sacB* (Schweizer, H. P., Klassen, T. and Hoang, T. (1996) Mol. Biol. of *Pseudomonas*. ASM press, 229-237 are available for *Pseudomonas*, *Salmonella* and *Klebsiella* species. *E. faecalis* genes can be disrupted by recombining in a non-replicating plasmid that contains an internal fragment to that gene (Leboeuf, C., L. Leblanc, Y. Auffray and A. Hartke. 2000. J. Bacteriol. 182:5799-5806).

The crossover PCR amplification product is subcloned into a suitable vector having a selectable marker, such as a drug resistance marker. In some embodiments the vector may have an origin of replication which is functional in *E. coli* or another organism distinct from the organism in which homologous recombination is to occur, allowing the plasmid to be grown in *E. coli* or the organism other than that in which homologous recombination is to occur, but may lack an origin of replication functional in *Staphylococcus aureus*, *Salmonella typhimurium*, *Klebsiella pneumoniae*, *Pseudomonas aeruginosa* and *Enterococcus faecalis*, *Escherichia coli*, *Enterococcus faecalis*, *Haemophilus influenzae*, *Helicobacter pylori*, *Klebsiella pneumoniae*, *Pseudomonas aeruginosa*, *Staphylococcus aureus*, or *Salmonella typhi* such that selection of the selectable marker requires integration of the vector into the homologous region of the *Staphylococcus aureus*, *Salmonella typhimurium*, *Klebsiella pneumoniae*, *Pseudomonas aeruginosa* and *Enterococcus faecalis*,

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Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa, Staphylococcus aureus, or Salmonella typhi chromosome. Usually a single crossover event is responsible for this integration event such that the Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa, Staphylococcus aureus, or Salmonella typhi chromosome now contains a tandem duplication of the target gene consisting of one wild type allele and one deletion null allele separated by vector sequence. Subsequent resolution of the duplication results in both removal of the vector sequence and either restoration of the wild type gene or replacement by the in-frame deletion. The latter outcome will not occur if the gene should prove essential. A more detailed description of this method is provided in Example 5 below. It will be appreciated that this method may be practiced with any of the nucleic acids or organisms described herein.

Recombinant DNA techniques can be used to express the entire coding sequences of the gene identified as required for proliferation, or portions thereof. The over-expressed proteins can be used as reagents for further study. The identified exogenous sequences are isolated, purified, and cloned into a suitable expression vector using methods well known in the art. If desired, the nucleic acids can contain the nucleotide sequences encoding a signal peptide to facilitate secretion of the expressed protein.

Expression of fragments of the bacterial genes identified as required for proliferation is also contemplated by the present invention. The fragments of the identified genes can encode a polypeptide comprising at least 5, at least 10, at least 15, at least 20, at least 25, at least 30, at least 35, at least 40, at least 45, at least 50, at least 55, at least 60, at least 65, at least 75, or more than 75 consecutive amino acids of a gene complementary to one of the identified sequences of the present invention. The nucleic acids inserted into the expression vectors can also contain endogenous sequences upstream and downstream of the coding sequence.

When expressing the encoded protien of the idnetified required for bacterial proliferation or a fragment thereof, the nucleotide sequence to be expressed is operably linked to a promoter in an expression vector using conventional cloning technology. The expression vector can be any of the bacterial, insect, yeast, or mammalian expression systems known in the art. Commercially available vectors and expression systems are available from a variety of suppliers including Genetics Institute (Cambridge, MA), Stratagene (La Jolla, California), Promega (Madison, Wisconsin), and Invitrogen (San Diego, California). If desired, to enhance expression and facilitate proper protein folding, the codon usage and codon bias of the sequence can be optimized for the particular expression organism in which the expression vector is introduced, as explained by Hatfield, et al., U.S. Patent No. 5,082,767. Fusion protein expression systems are also contemplated by the present invention.

Following expression of the protein encoded by the identified exogenous nucleic acid, the protein may be purified. Protein purification techniques are well known in the art. Proteins encoded and expressed from identified exogenous nucleic acids can be partially purified using precipitation techniques, such as precipitation with polyethylene glycol. Alternatively, epitope tagging of the protein can be used to allow simple one step purification of the protein. In addition, chromatographic methods such as ion-exchange chromatography, gel filtration, use of hydroxyapaptite columns, immobilized reactive dyes, chromatofocusing, and use of high-performance liquid chromatography, may also be used to purify the protein. Electrophoretic methods such as one-dimensional gel electrophoresis, high-resolution two-dimensional polyacrylamide electrophoresis, isoelectric focusing, and others are contemplated as purification methods. Also, affinity chromatographic methods, comprising antibody columns, ligand presenting columns and other affinity chromatographic matrices are contemplated as purification methods in the present invention.

The purified proteins produced from the gene coding sequences identified as required for proliferation can be used in a variety of protocols to generate useful antimicrobial reagents. In one embodiment of the present invention, antibodies are generated against the proteins expressed from the identified exogenous nucleic acids. Both monoclonal and polyclonal antibodies can be generated against the expressed proteins. Methods for generating monoclonal and polyclonal antibodies are well known in the art. Also, antibody fragment preparations prepared from the produced antibodies discussed above are contemplated.

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In addition, the purified protein, fragments thereof, or derivatives thereof may be administered to an individual in a pharmaceutically acceptable carrier to induce an immune response against the protein. Preferably, the immune response is a protective immune response which protects the individual. Methods for determining appropriate dosages of the protein and pharmaceutically acceptable carriers may be determined empiracally and are familiar to those skilled in the art.

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Another application for the purified proteins of the present invention is to screen small molecule libraries for candidate compounds active against the various target proteins of the present invention. Advances in the field of combinatorial chemistry provide methods, well known in the art, to produce large numbers of candidate compounds that can have a binding, or otherwise inhibitory effect on a target protein. Accordingly, the screening of small molecule libraries for compounds with binding affinity or inhibitory activity for a target protein produced from an identified gene is contemplated by the present invention.

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The present invention further contemplates utility against a variety of other pathogenic microorganisms in addition to Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa, Staphylococcus aureus, or Salmonella typhi. For example, homologous coding nucleic acids, homologous antisense nucleic acids or homologous polypeptides from other pathogenic

microorganisms (including nucleic acids homologous to the nucleic acids of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, nucleic acids homologous to the antisense nucleic acids of SEQ ID NOs.: 8-3795, and polypeptides homologous to the polypeptides of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110) may be identified using methods such as those described herein. The homologous coding nucleic acids, homologous antisense nucleic acids or homologous polypeptides may be used to identify compounds which inhibit the proliferation of these other pathogenic microorganisms using methods such as those described herein.

For example, the proliferation-required nucleic acids, antisense nucleic acids, and polypeptides from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae. Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa, Staphylococcus aureus, or Salmonella typhi described herein (including the nucleic acids of SEO ID NOs.: 3796-3800, 3806-4860, 5916-10012, the antisense nucleic acids of SEO ID NOs; 8-3795, and the polypeptides of SEO ID NOs.: 3801-3805, 4861-5915, 10013-14110) may be used to identify homologous coding nucleic acids, homologous antisense nucleic acids or homologous polypeptides required for proliferation in prokaryotes and eukaryotes. For example, nucleic acids or polypeptides required for the proliferation of protists, such as *Plasmodium* spp.; plants; animals, such as Entamoeba spp. and Contracaecum spp; and fungi including Candida spp., (e.g., Candida albicans), Cryptococcus neoformans, and Aspergillus fumigatus may be identified. In one embodiment of the present invention, monera, specifically bacteria, including both Gram positive and Gram negative bacteria, are probed in search of novel gene sequences required for proliferation. Likewise. homologous antisense nucleic acids which may be used to inhibit growth of these organisms or to identify antibiotics may also be identified. These embodiments are particularly important given the rise of drug resistant bacteria.

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The number of bacterial species that are becoming resistant to existing antibiotics is growing. A partial list of these microorganisms includes: Escherichia spp., such as E. coli, Enterococcus spp, such as E. faecalis; Pseudomonas spp., such as P. aeruginosa, Clostridium spp., such as C. botulinum, Haemophilus spp., such as H. influenzae, Enterobacter spp., such as E. cloacae, Vibrio spp., such as V. cholera; Moraxala spp., such as M. catarrhalis; Streptococcus spp., such as S. pneumoniae, Neisseria spp., such as N. gonorrhoeae; Mycoplasma spp., such as Mycoplasma pneumoniae; Salmonella typhimurium; Helicobacter pylori; Escherichia coli; and Mycobacterium tuberculosis. The genes and polypeptides identified as required for the proliferation of Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa, Staphylococcus aureus, or Salmonella typhi (including the nucleic acids of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, the sequences complementary to the nucleic acids of SEQ ID NOs.: 3796-3800, 3806-4860,

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5916-10012, and the polypeptides of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110) can be used to identify homologous coding nucleic acids or homologous polypeptides required for proliferation from these and other organisms using methods such as nucleic acid hybridization and computer database analysis. Likewise, the antisense nucleic acids which inhibit proliferation of Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa, Staphylococcus aureus, or Salmonella typhi (including the antisense nucleic acids of SEQ ID NOs.: 8-3795 or the sequences complementary thereto) may also be used to identify antisense nucleic acids which inhibit proliferation of these and other microorganisms or cells using nucleic acid hybridization or computer database analysis.

In one embodiment of the present invention, the nucleic acid sequences from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae. 15 Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa, Staphylococcus aureus, or Salmonella typhii (including the nucleic acids of SEO ID NOs.: 3796-3800, 3806-4860, 5916-10012 and the antisense nucleic acids of SEQ ID NOs. 8-3795) are used to screen genomic libraries generated from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, 20 Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa. Staphylococcus aureus, or Salmonella typhi and other bacterial species of interest. For example, the genomic library may be from Gram positive bacteria, Gram negative bacteria or other organisms including Anaplasma marginale, Aspergillus funigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida 25 glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, 30 Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae. Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella 35 typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica,

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Yersinia pestis or any species falling within the genera of any of the above species, including coagulase negative species of Staphylococcus. In some embodiments, the genomic library may be from an organism other than E. coli. Standard molecular biology techniques are used to generate genomic libraries from various cells or microorganisms. In one aspect, the libraries are generated and bound to nitrocellulose paper. The identified exogenous nucleic acid sequences of the present invention can then be used as probes to screen the libraries for homologous sequences.

For example, the libraries may be screened to identify homologous coding nucleic acids or homologous antisense nucleic acids comprising nucleotide sequences which hybridize under stringent conditions to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795. nucleic acids comprising nucleotide sequences which hybridize under stringent conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of one of SEO ID NOs, 8-3795, nucleic acids comprising nucleotide sequences which hybridize under stringent conditions to a nucleic acid complementary to one of SEO ID NOs. 8-3795, nucleic acids comprising nucleotide sequences which hybridize under stringent conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200. 300, 400, or 500 consecutive nucleotides of the sequence complementary to one of SEO ID NOs. 8-3795, nucleic acids comprising nucleotide sequences which hybridize under stringent conditions to a nucleic acid selected from the group consisting of SEO ID NOS.: 3796-3800, 3806-4860, 5916-10012, nucleic acids comprising nucleotide sequences which hybridize under stringent conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of one of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, nucleic acids comprising nucleotide sequences which hybridize under stringent conditions to a nucleic acid complementary to one of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, nucleic acids comprising nucleotide sequences which hybridize under stringent conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of the sequence complementary to one of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, nucleic acids comprising nucleotide sequences which hybridize under stringent conditions to a nucleic acid selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, and nucleic acids comprising nucleotide sequences which hybridize under stringent conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of one of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012.

The libraries may also be screened to identify homologous nucleic coding nucleic acids or homologous antisense nucleic acids comprising nucleotide sequences which hybridize under moderate conditions to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795, nucleic acids comprising nucleotide sequences which hybridize under moderate conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of one of SEQ ID NOs. 8-3795, nucleic acids comprising nucleotide

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sequences which hybridize under moderate conditions to a nucleic acid complementary to one of SEQ ID NOs. 8-3795, nucleic acids comprising nucleotide sequences which hybridize under moderate conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of the sequence complementary to one of SEQ ID NOs. 8-3795, nucleic acids comprising nucleotide sequences which hybridize under moderate conditions to a nucleic acid selected from the group consisting of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, nucleic acids comprising nucleic acid sequences which hybridize under moderate conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of one of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, nucleic acids comprising nucleotide sequences which hybridize under moderate conditions to a nucleic acid complementary to one of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012 and nucleic acids comprising nucleotide sequences which hybridize under moderate conditions to a fragment comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of the sequence complementary to one of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012.

The homologous nucleic coding nucleic acids, homologous antisense nucleic acids or homologous polypeptides identified as above can then be used as targets or tools for the identification of new, antimicrobial compounds using methods such as those described herein. In some embodiments, the homologous coding nucleic acids, homologous antisense nucleic acids, or homologous polypeptides may be used to identify compounds with activity against more than one microorganism.

For example, the preceding methods may be used to isolate homologous coding nucleic acids or homologous antisense nucleic acids comprising a nucleotide sequence with at least 97%, at least 95%, at least 90%, at least 85%, at least 80%, or at least 70% nucleotide sequence identity to a nucleotide sequence selected from the group consisting of one of the sequences of SEO ID NOS. 8-3795, fragments comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides thereof, and the sequences complementary thereto. The preceding methods may also be used to isolate homologous coding nucleic acids or homologous antisense nucleic acids comprising a nucleotide sequence with at least 97%, at least 95%, at least 90%, at least 85%, at least 80%, or at least 70% nucleotide sequence identity to a nucleotide sequence selected from the group consisting of one of the nucleotide sequences of SEO ID NOS.: 3796-3800, 3806-4860. 5916-10012, fragments comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides thereof, and the sequences complementary thereto. In some embodiments, the preceding methods may be used to isolate homologous coding nucleic acids or homologous antisense nucleic acids comprising a nucleotide sequence with at least 97%, at least 95%, at least 90%, at least 85%, at least 80%, or at least 70% nucleotide sequence identity to a nucleic acid sequence selected from the group consisting of one of the sequences of SEO ID NOS.

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3796-3800, 3806-4860, 5916-10012, fragments comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides thereof, and the sequences complementary thereto. Identity may be measured using BLASTN version 2.0 with the default parameters. (Altschul, S.F. et al. Gapped BLAST and PSI-BLAST: A New Generation of Protein Database Search Programs, Nucleic Acid Res. 25: 3389-3402 (1997)). For example, the homologous polynucleotides may comprise a coding sequence which is a naturally occurring allelic variant of one of the coding sequences described herein. Such allelic variants may have a substitution, deletion or addition of one or more nucleotides when compared to the nucleic acids of SEQ ID NOs: 8-3795, SEQ ID NOS:: 3796-3800, 3806-4860, 5916-10012 or the nucleotide sequences complementary thereto.

Additionally, the above procedures may be used to isolate homologous coding nucleic acids which encode polypeptides having at least 99%, 95%, at least 90%, at least 85%, at least 80%, at least 70%, at least 60%, at least 50%, at least 40% or at least 25% amino acid identity or similarity to a polypeptide comprising the sequence of one of SEQ ID NOs: 3801-3805, 4861-5915, 10013-14110 or to a polypeptide whose expression is inhibited by a nucleic acid of one of SEQ ID NOs: 8-3795 or fragments comprising at least 5, 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, or 150 consecutive amino acids thereof as determined using the FASTA version 3.0t78 algorithm with the default parameters. Alternatively, protein identity or similarity may be identified using BLASTP with the default parameters, BLASTX with the default parameters, or TBLASTN with the default parameters. (Altschul, S.F. et al. Gapped BLAST and PSI-BLAST: A New Generation of Protein Database Search Programs, Nucleic Acid Res. 25: 3389-3402 (1997)).

Alternatively, homologous coding nucleic acids, homologous antisense nucleic acids or homologous polypeptides may be identified by searching a database to identify sequences having a desired level of nucleotide or amino acid sequence homology to a nucleic acid or polypertide involved in proliferation or an antisense nucleic acid to a nucleic acid involved in microbial proliferation. A variety of such databases are available to those skilled in the art, including GenBank and GenSeq. In some embodiments, the databases are screened to identify nucleic acids with at least 97%, at least 95%, at least 90%, at least 85%, at least 80%, or at least 70% nucleotide sequence identity to a nucleic acid required for proliferation, an antisense nucleic acid which inhibits proliferation, or a portion of a nucleic acid required for proliferation or a portion of an antisense nucleic acid which inhibits proliferation. For example, homologous coding sequences may be identified by using a database to identify nucleic acids homologous to one of SEO ID Nos. 8-3795, homologous to fragments comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides thereof, nucleic acids homologous to one of SEQ ID NOS.: 3796-3800, 3806-4860, 5916-10012, homologous to fragments comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200, 300, 400, or 500 consecutive nucleotides of one of SEO ID NOS.: 3796-3800, 3806-4860, 5916-10012, nucleic acids homologous to one of SEO ID Nos. 8-

3795, homologous to fragments comprising at least 10, 15, 20, 25, 30, 35, 40, 50, 75, 100, 150, 200. 300, 400, or 500 consecutive nucleotides thereof or nucleic acids homologous to the sequences complementary to any of the preceding nucleic acids. In other embodiments, the databases are screened to identify polypeptides having at least 99%, 95%, at least 90%, at least 85%, at least 80%, 5 at least 70%, at least 60%, at least 50%, at least 40% or at least 25% amino acid sequence identity or similarity to a polypeptide involved in proliferation or a portion thereof. For example, the database may be screened to identify polypeptides homologous to a polypeptide comprising one of SEQ ID NOs: 3801-3805, 4861-5915, 10013-14110, a polypeptide whose expression is inhibited by a nucleic acid of one of SEO ID NOs: 8-3795 or homologous to fragments comprising at least 5, 10. 10 15, 20, 25, 30, 35, 40, 50, 75, 100, or 150 consecutive amino acids of any of the preceding polypeptides. In some embodiments, the database may be screened to identify homologous coding nucleic acids, homologous antisense nucleic acids or homologous polypeptides from cells or microorganisms other than the Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus 15 faecalis, Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa, Staphylococcus aureus, or Salmonella typhi species from which they were obtained. For example the database may be screened to identify homologous coding nucleic acids, homologous antisense nucleic acids or homologous polypeptides from microorganisms such as Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella 20 pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae. Chlamydia trachomatus, Clostridium botulinum. Clostridium difficile. Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus 25 neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, 30 Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis or any species falling within the genera of any of the above species, including 35 coagulase negative Staphylococcus. In some embodiments, the homologous coding nucleic acids. homologous antisense nucleic acids, or homologous polypeptides are from an organism other than E. coli.

In another embodiment, gene expression arrays and microarrays can be employed. Gene expression arrays are high density arrays of DNA samples deposited at specific locations on a glass chip, nylon membrane, or the like. Such arrays can be used by researchers to quantify relative gene expression under different conditions. Gene expression arrays are used by researchers to help identify optimal drug targets, profile new compounds, and determine disease pathways. An example of this technology is found in U.S. Patent No. 5807522.

It is possible to study the expression of all genes in the genome of a particular microbial organism using a single array. For example, the arrays may consist of 12 x 24 cm nylon filters containing PCR products corresponding to ORFs from *Staphylococcus aureus*, *Salmonella typhimurium*, *Klebsiella pneumoniae*, *Pseudomonas aeruginosa* and *Enterococcus faecalis*, *Escherichia coli*, *Enterococcus faecalis*, *Haemophilus influenzae*, *Helicobacter pylori*, *Klebsiella pneumoniae*, *Pseudomonas aeruginosa*, *Staphylococcus aureus*, or *Salmonella typhi* (including the nucleic acids of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012). 10 ngs of each PCR product are spotted every 1.5 mm on the filter. Single stranded labeled cDNAs are prepared for hybridization to the array (no second strand synthesis or amplification step is done) and placed in contact with the filter. Thus the labeled cDNAs are of "antisense" orientation. Quantitative analysis is done by phosphorimager.

Hybridization of cDNA made from a sample of total cell mRNA to such an array followed by detection of binding by one or more of various techniques known to those in the art results in a signal at each location on the array to which cDNA hybridized. The intensity of the hybridization signal obtained at each location in the array thus reflects the amount of mRNA for that specific gene that was present in the sample. Comparing the results obtained for mRNA isolated from cells grown under different conditions thus allows for a comparison of the relative amount of expression of each individual gene during growth under the different conditions.

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Gene expression arrays may be used to analyze the total mRNA expression pattern at various time points after induction of an antisense nucleic acid complementary to a proliferation-required gene. Analysis of the expression pattern indicated by hybridization to the array provides information on other genes whose expression is influenced by antisense expression. For example, if the antisense is complementary to a gene for ribosomal protein L7/L12 in the 50S subunit, levels of other mRNAs may be observed to increase, decrease or stay the same following expression of antisense to the L7/L12 gene. If the antisense is complementary to a different 50S subunit ribosomal protein mRNA (e.g. L25), a different mRNA expression pattern may result. Thus, the mRNA expression pattern observed following expression of an antisense nucleic acid comprising a nucleotide sequence complementary to a proliferation required gene may identify other proliferation-required nucleic acids. In addition, the mRNA expression patterns observed when the bacteria are exposed to candidate drug compounds or known antibiotics may be compared to those observed with antisense nucleic acids comprising a nucleotide sequence complementary to a

proliferation-required nucleic acid. If the mRNA expression pattern observed with the candidate drug compound is similar to that observed with the antisense nucleic acid, the drug compound may be a promising therapeutic candidate. Thus, the assay would be useful in assisting in the selection of promising candidate drug compounds for use in drug development.

In cases where the source of nucleic acid deposited on the array and the source of the nucleic acid being hybridized to the array are from two different cells or microorganisms, gene expression arrays can identify homologous nucleic acids in the two cells or microorganisms.

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The present invention also contemplates additional methods for screening other microorganisms for proliferation-required genes. In one aspect of this embodiment, an antisense nucleic acid comprising a nucleotide sequence complementary to the proliferation-required sequences from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa, Staphylococus aureus, or Salmonella typhi or a portion thereof is transcribed in an antisense orientation in such a way as to alter the level or activity of a nucleic acid required for proliferation of an autologous or heterologous cell or microorganism. For example, the antisense nucleic acid may be a homologous antisense nucleic acid such as an antisense nucleic acid homologous to the nucleotide sequence complementary to one of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, an antisense nucleic acid comprising a nucleotide sequence homologous to one of SEO ID Nos.; 8-3795, or an antisense nucleic acid comprising a nucleotide sequence complementary to a portion of any of the preceding nucleic acids. The cell or microorganism transcribing the homologous antisense nucleic acid may be used in a cell-based assay, such as those described herein, to identify candidate antibiotic compounds. In another embodiment, the conserved portions of nucleotide sequences identified as proliferationrequired can be used to generate degenerate primers for use in the polymerase chain reaction (PCR). The PCR technique is well known in the art. The successful production of a PCR product using degenerate probes generated from the nucleotide sequences identified herein indicates the presence of a homologous gene sequence in the species being screened. This homologous gene is then isolated, expressed, and used as a target for candidate antibiotic compounds. In another aspect of this embodiment, the homologous gene (for example a homologous coding nucleic acid)thus identified, or a portion thereof, is transcribed in an autologous cell or microorganism or in a heterologous cell or microorganism in an antisense orientation in such a way as to alter the level or activity of a homologous gene required for proliferation in the autologous or heterologous cell or microorganism. Alternatively, a homologous antisense nucleic acid may be transcribed in an autologous or heterologous cell or microorganism in such a way as to alter the level or activity of a gene product required for proliferation in the autologous or heterologous cell or microorganism.

The nucleic acids homologous to the genes required for the proliferation of Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and

Enterococcus faecalis. Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa, Staphylococcus aureus, or Salmonella typhi or the sequences complementary thereto may be used to identify homologous coding nucleic acids or homologous antisense nucleic acids from cells or microorganisms other than Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa 5 and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae. Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa, Staphylococcus aureus, or Salmonella typhi to inhibit the proliferation of cells or microorganisms other than Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, 10 Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa, Staphylococcus aureus, or Salmonella typhi by inhibiting the activity or reducing the amount of the identified homologous coding nucleic acid or homologous polypeptide in the cell or microorganism other than Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa 15 Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi or to identify compounds which inhibit the growth of cells or microorganisms other than Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi as described below. For example, the nucleic acids homologous to proliferation-required genes from Staphylococcus aureus, 20 Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa, Staphylococcus aureus, or Salmonella typhi or the sequences complementary thereto may be used to identify compounds which inhibit the growth 25 of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus 30 neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella 35 multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella

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boydii, Shigella dysenteriae, Shigella flexneri, Shigella somnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the genera of any of the above species. In some embodiments of the present invention, the nucleic acids homologous to proliferation-required sequences from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa, Staphylococcus aureus, or Salmonella typhi (including nucleic acids homologous to one of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012) or the sequences complementary thereto (including nucleic acids homologous to one of SEQ ID NOs.: 8-3795) are used to identify proliferation-required sequences in an organism other than E. coli.

In another embodiment of the present invention, antisense nucleic acids complementary to the

sequences identified as required for proliferation or portions thereof (including antisense nucleic acids comprising a nucleotide sequence complementary to one of SEQ ID NOs.: 3796-3800, 3806-4860, 15 5916-10012 or portions thereof, such as the nucleic acids of SEQ ID NOs.: 8-3795) are transferred to vectors capable of function within a species other than the species from which the sequences were obtained. For example, the vector may be functional in Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia. Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), 20 Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corvuebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae. Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, 25 Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes. Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, 30 Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis or any species falling within the genera of any of the above species. In some embodiments of the present invention, the vector may be functional in an organism other than E. coli. As would be 35 appreciated by one of ordinary skill in the art, vectors may contain certain elements that are species specific. These elements can include promoter sequences, operator sequences, repressor genes, origins of replication, ribosomal binding sequences, termination sequences, and others. To use the

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antisense nucleic acids, one of ordinary skill in the art would know to use standard molecular biology techniques to isolate vectors containing the sequences of interest from cultured bacterial cells, isolate and purify those sequences, and subclone those sequences into a vector adapted for use in the species of bacteria to be screened.

Vectors for a variety of other species are known in the art. For example, numerous vectors which function in *E. coli* are known in the art. Also, Pla et al. have reported an expression vector that is functional in a number of relevant hosts including: *Salmonella typhimurium, Pseudomonas putida, and Pseudomonas aeruginosa*. J. Bacteriol. 172(8):4448-55 (1990). Brunschwig and Darzins (Gene (1992) 111:35-4) described a shuttle expression vector for *Pseudomonas aeruginosa*. Similarly many examples exist of expression vectors that are freely transferable among various Gram-positive microorganisms. Expression vectors for *Enterococcus faecalis* may be engineered by incorporating suitable promoters into a pAK80 backbone (Israelsen, H., S. M. Madsen, A. Vrang, E. B. Hansen and E. Johansen. 1995. Appl. Environ. Microbiol. 61:2540-2547).

Following the subcloning of the antisense nucleic acids complementary to proliferationrequired sequences from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae, Pseudomonas aeruginosa. Staphylococcus aureus, or Salmonella typhi or portions thereof into a vector functional in a second cell or microorganism of interest (i.e. a cell or microorganism other than the one from which the identified nucleic acids were obtained), the antisense nucleic acids are conditionally transcribed to test for bacterial growth inhibition. The nucleotide sequences of the nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi that, when transcribed, inhibit growth of the second cell or microorganism are compared to the known genomic sequence of the second cell or microorganism to identify the homologous gene from the second organism. If the homologous sequence from the second cell or microorganism is not known, it may be identified and isolated by hybridization to the proliferation-required Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi sequence of interest or by amplification using PCR primers based on the proliferation-required nucleotide sequence of interest as described above. In this way, sequences which may be required for the proliferation of the second cell or microorganism may be identified. For example, the second microorganism may be Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis,

Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinti, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis or any species falling within the genera of any of the above species. In some embodiments of the present invention, the second microorganism is an organism other than E. coli.

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The homologous nucleic acid sequences from the second cell or microorganism which are 15 identified as described above may then be operably linked to a promoter, such as an inducible promoter, in an antisense orientation and introduced into the second cell or microorganism. The techniques described herein for identifying Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa and Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, Klebsiella pneumoniae, 20 Pseudomonas aeruginosa, Staphylococcus aureus, or Salmonella typhi genes required for proliferation may thus be employed to determine whether the identified nucleotide sequences from a second cell or microorganism inhibit the proliferation of the second cell or microorganism. For example, the second microorganism may be Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, 25 Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus 30 faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus yulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, 35 Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans,

Treponema pallidum, Yersinia enterocolitica, Yersinia pestis or any species falling within the genera of any of the above species. In some embodiments of the present invention, the second microorganism may be an organism other than E. coli.

Antisense nucleic acids required for the proliferation of microorganisms other than 5 Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi or the genes corresponding thereto, may also be hybridized to a microarray containing the Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis ORFs, Escherichia coli. 10 Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, and Salmonella typhi (including the nucleic acids of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012) to gauge the homology between the Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi sequences and the proliferation-15 required nucleic acids from other cells or microorganisms. For example, the proliferation-required nucleic acid may be from Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida 20 pseudotropicalis), Candida dubliniensis, Chlamvdia pneumoniae, Chlamvdia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis. Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, 25 Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, 30 Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis or any species falling within the genera of any of the above species. In some embodiments of the present invention, the proliferation-required nucleotide sequences from Staphylococcus aureus, Salmonella typhimurium. Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli. 35 Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, Salmonella typhi or homologous nucleic acids are used to identify proliferation-required sequences in an organism other than E. coli. In some embodiments of the present invention, the proliferation-required sequences

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may be from an organism other than E. coli. The proliferation-required nucleic acids from a cell or microorganism other than Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi may be hybridized to the array under a variety of conditions which permit hybridization to occur when the probe has different levels of homology to the nucleotide sequence on the microarray. This would provide an indication of homology across the cells or microorganisms as well as clues to other possible essential genes in these cells or microorganisms.

In still another embodiment, the antisense nucleic acids of the present invention (including the antisense nucleic acids of SEQ ID NOs. 8-3795 or homologous antisense nucleic acids) that inhibit bacterial growth or proliferation can be used as antisense therapeutics for killing bacteria. The antisense sequences can be complementary to one of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, homologous nucleic acids, or portions thereof. Alternatively, antisense therapeutics can be complementary to operons in which proliferation-required genes reside (i.e. the antisense nucleic acid may hybridize to a nucleotide sequence of any gene in the operon in which the proliferation-required genes reside). Further, antisense therapeutics can be complementary to a proliferation-required gene or portion thereof with or without adjacent noncoding sequences, an intragenic sequence (i.e. a sequence within a gene), an intergenic sequence (i.e. a sequence between genes), a sequence spanning at least a portion of two or more genes, a 5' noncoding region or a 3' noncoding region located upstream or downstream from the actual sequence that is required for bacterial proliferation or an operon containing a proliferation-required gene.

In addition to therapeutic applications, the present invention encompasses the use of nucleic acids complementary to nucleic acids required for proliferation as diagnostic tools. For example, nucleic acid probes comprising nucleotide sequences complementary to proliferation-required sequences that are specific for particular species of cells or microorganisms can be used as probes to identify particular microorganism species or cells in clinical specimens. This utility provides a rapid and dependable method by which to identify the causative agent or agents of a bacterial infection. This utility would provide clinicians the ability to accurately identify the species responsible for the infection and amdminister a compound effective against it. In an extension of this utility, antibodies generated against proteins translated from mRNA transcribed from proliferation-required sequences can also be used to screen for specific cells or microorganisms that produce such proteins in a species-specific manner.

Other embodiments of the present invention include methods of identifying compounds which inhibit the activity of gene products required for cellular proliferation using rational drug design. As discussed in more detail below, in such methods, the structure of the gene product is determined using techniques such as x-ray crystallography or computer modeling. Compounds are screened to identify those which have a structure which would allow them to interact with the gene product or a portion

thereof to inhibit its activity. The compounds may be obtained using any of a variety of methods familiar to those skilled in the art, including combinatorial chemistry. In some embodiments, the compounds may be obtained from a natural product library. In some embodiments, compounds having a structure which allows them to interact with the active site of a gene product, such as the active site of an enzyme, or with a portion of the gene product which interacts with another biomolecule to form a complex are identified. If desired, lead compounds may be identified and further optimized to provide compounds which are highly effective against the gene product.

The following examples teach the genes of the present invention and a subset of uses for the genes identified as required for proliferation. These examples are illustrative only and are not intended to limit the scope of the present invention.

EXAMPLES

The following examples are directed to the identification and exploitation of genes required for proliferation. Methods of gene identification are discussed as well as a variety of methods to utilize the identified sequences. It will be appreciated that any of the antisense nucleic acids, proliferartion-required genes or proliferation-required gene products described herein, or portions thereof, may be used in the procedures described below, including the antisense nucleic acids of SEQ ID NOs.: 8-3795, the nucleic acids of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, or the polypeptides of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110. Likewise, homologous coding nucleic acids or portions thereof, may be used in any of the procedures described below.

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Genes Identified as Required for Proliferation of Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa or Enterococcus faecalis

Genomic fragments were operably linked to an inducible promoter in a vector and assayed for growth inhibition activity. Example 1 describes the examination of a library of genomic fragments cloned into vectors comprising inducible promoters. Upon induction with xylose or IPTG, the vectors produced an RNA molecule corresponding to the subcloned genomic fragments. In those instances where the genomic fragments were in an antisense orientation with respect to the promoter, the transcript produced was complementary to at least a portion of an mRNA (messenger RNA) encoding a Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa or Enterococcus faecalis gene product such that they interacted with sense mRNA produced from various Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa or Enterococcus faecalis genes and thereby decreased the translation efficiency or the level of the sense messenger RNA thus decreasing production of the protein encoded by these sense mRNA molecules. In cases where the sense mRNA encoded a protein required for proliferation, bacterial cells containing a vector from which transcription from the promoter had been induced failed to grow or grew at a substantially reduced rate. Additionally, in cases where the transcript produced was complementary to at least a portion of a non-translated RNA and where that

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non-translated RNA was required for proliferation, bacterial cells containing a vector from which transcription from the promoter had been induced also failed to grow or grew at a substantially reduced rate.

EXAMPLE 1

Inhibition of Bacterial Proliferation after Induction of Antisense Expression

Nucleic acids involved in proliferation of Staphylococcus aureus, Salmonella typhimurium, and Klebsiella pneumoniae were identified as follows. Randomly generated fragments of Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa or Enterococcus faecalis genomic DNA were transcribed from inducible promoters.

In the case of Staphylococcus aureus, a novel inducible promoter system, XyIT5, comprising a modified T5 promoter fused to the xylO operator from the xylA promoter of Staphylococcus aureus was used. The promoter is described in U.S. Provisional Patent Application Serial Number 60/259,434. Transcription from this hybrid promoter is inducible by xylose.

Randomly generated fragments of Salmonella typhimurium genomic DNA were transcribed from an IPTG inducible promoter in pLEX5BA (Krause et al., J. Mol. Biol. 274: 365 (1997) or a derivative thereof. Randomly generated fragements of Klebsiella pneumoniae genomic DNA were expressed from an IPTG inducible promoter in pLEX5BA-Kan. To construct pLEX5BA-kan, pLEX5BA was digested to completion with ClaI in order to remove the bla gene. Then the plasmid was treated with a partial NotI digestion and blunted with T4 DNA polymerase. A 3.2 kbp fragment was then gel purified and ligated to a blunted 1.3 kbp kan gene from pKanπ. Kan resistant transformants were selected on Kan plates. Orientation of the kan gene was checked by SmaI digestion. A clone, which had the kan gene in the same orientation as the bla gene, was used to identify genes required for proliferation of Klebsiella pneumoniae.

Randomly generated fragments of *Pseudomonas aeruginosa* genomic DNA were trancribed from a two-component inducible promoter system. Integrated on the chromosome was the T7 RNA polymerase gene regulated by *lacUV5/lacO* (Brunschwig, E. and Darzins, A. 1992. Gene 111:35-41). On a separate plasmid, a T7 gene 10 promoter, which is transcribed by T7 RNA polymerase, was fused with a *lacO* operator followed by a multiple cloning site.

Should the genomic DNA downstream of the promoter contain, in an antisense orientation, at least a portion of an mRNA or a non-translated RNA encoding a gene product involved in proliferation, then induction of transcription from the promoter will result in detectable inhibition of proliferation.

In the case of *Staphylococcus aureus*, a shotgun library of *Staphylococcus aureus* genomic fragments was cloned into the vector pXyIT5-P15a, which harbors the XyIT5 inducible promoter. The vector was linearized at a unique *Bam*HI site immediately downstream of the XyIT5 promoter/operator. The linearized vector was treated with shrimp alkaline phosphatase to prevent reclosure of the linearized ends. Genomic DNA isolated from *Staphylococcus aureus* strain RN450

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was fully digested with the restriction enzyme Sau3A, or, alternatively, partially digested with DNase I and "blunt-ended" by incubating with T4 DNA polymerase. Random genomic fragments between 200 and 800 base pairs in length were selected by gel purification. The size-selected genomic fragments were added to the linearized and dephosphorylated vector at a molar ratio of 0.1 to 1, and ligated to form a shotgun library.

The ligated products were transformed into electrocompetent *E. coli* strain XL1-Blue MRF (Stratagene) and plated on LB medium with supplemented with carbenicillin at 100 μ g/ml. Resulting colonies numbering 5 x 10⁵ or greater were scraped and combined, and were then subjected to plasmid purification.

The purified library was then transformed into electrocompetent *Staphylococcus aureus* RN4220. Resulting transformants were plated on agar containing LB + 0.2% glucose (LBG medium) + chloramphenicol at 15 µg/ml (LBG+CM15 medium) in order to generate 100 to 150 platings at 500 colonies per plating. The colonies were subjected to robotic picking and arrayed into wells of 384 well culture dishes. Each well contained 100µl of LBG + CM15 liquid medium. Inoculated 384 well dishes were incubated 16 hours at 37°C, and each well was robotically gridded onto solid LBG + CM15 medium with or without 2% xylose. Gridded plates were incubated 16 hours at 37°C, and then manually scored for arrayed colonies that were growth-compromised in the presence of xylose.

Arrayed colonies that were growth-sensitive on medium containing 2% xylose, yet were able to grow on similar medium lacking xylose, were subjected to further growth sensitivity analysis as follows: Colonies from the plate lacking xylose were manually picked and inoculated into individual wells of a 96 well culture dish containing LBG + CM15, and were incubated for 16 hours at 37°C. These cultures were robotically diluted 1/100 into fresh medium and allowed to incubate for 4 hours at 37°C, after which they were subjected to serial dilutions in a 384 well array and then gridded onto media containing 2% xylose or media lacking xylose. After growth for 16 hours at 37°C, the arrays that resulted on the two media were compared to each other. Clones that grew similarly at all dilutions on both media were scored as a negative and were no longer considered. Clones that grew on xylose medium but failed to grow at the same serial dilution on the non-xylose plate were given a score based on the differential, i.e. should the clone grow at a serial dilution of 10⁴ or less on the xylose plate and grow at a serial dilution of 10⁸ or less on the non-xylose plate, then the corresponding clone received a score of "4" representing the log difference in growth observed.

For Salmonella typhimurium and Klebsiella pneumoniae growth curves were carried out by back diluting cultures 1:200 into fresh media containing 1 mM IPTG or media lacking IPTG and measuring the OD₄₅₀ every 30 minutes (min). To study the effects of transcriptional induction on solid medium, 10^2 , 10^3 , 10^4 , 10^5 , 10^6 , 10^7 and 10^8 fold dilutions of overnight cultures were prepared.

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Aliquots of from 0.5 to 3 μ l of these dilutions were spotted on selective agar plates with or without 1 mM IPTG. After overnight incubation, the plates were compared to assess the sensitivity of the clones to IPTG.

Nucleic acids involved in proliferation of *Pseudomonas aeruginosa* were identified as follows. Randomly generated fragments of *Pseudomonas aeruginosa* genomic DNA were transcribed from a two-component inducible promoter system. Integrated on the chromosome was the T7 RNA polymerase gene regulated by *lac*UV5/ *lac*O (Brunschwig, E. and Darzins, A. 1992. Gene 111:35-41). On an expression plasmid there was a T7 gene 10 promoter, which is transcribed by T7 RNA polymerase, fused with a *lac*O operator followed by a multiple cloning site.

Transcription from this hybrid promoter is inducible by IPTG. Should the genomic DNA downstream of the promoter contain, in an antisense orientation, at least a portion of an mRNA encoding a gene product involved in proliferation, then induction of expression from the promoter will result in detectable inhibition of proliferation.

A shotgun library of *Pseudomonas aeruginosa* genomic fragments was cloned into the vectors pEP5, pEP5S, or other similarly constructed vectors which harbor the T7lacO inducible promoter. The vector was linearized at a unique *Sma*I site immediately downstream of the T7lacO promoter/operator. The linearized vector was treated with shrimp alkaline phosphatase to prevent reclosure of the linearized ends. Genomic DNA isolated from *Pseudomonas aeruginosa* strain PAO1 was partially digested with DNase I and "blunt-ended" by incubating with T4 DNA polymerase. Random genomic fragments between 200 and 800 base pairs in length were selected by gel purification. The size-selected genomic fragments were added to the linearized and dephosphorylated vector at a molar ratio of 2 to 1, and ligated to form a shotgun library.

The ligated products were transformed into electrocompetent *E. coli* strain XL1-Blue MRF (Stratagene) and plated on LB medium with carbenicillin at 100 g/ml or Streptomycin 100 g/ml. Resulting colonies numbering 5 x 10⁵ or greater were scraped and combined, and were then subjected to plasmid purification.

The purified library was then transformed into electrocompetent *Pseudomonas aeruginosa* strain PAO1. Resulting transformants were plated on LB agar with carbenicillin at 100 g/ml or Streptomycin 40 g/ml in order to generate 100 to 150 platings at 500 colonies per plating. The colonies were subjected to robotic picking and arrayed into wells of 384 well culture dishes. Each well contained 100 1 of LB + CB 100 or Streptomycin 40 liquid medium. Inoculated 384 well dishes were incubated 16 hours at room temperature, and each well was robotically gridded onto solid LB + CB100 or Streptomycin 40 medium with or without 1 mM IPTG. Gridded plates were incubated 16 hours at 37°C, and then manually scored for arrayed colonies that were growth-compromised in the presence of IPTG.

Arrayed colonies that were growth-sensitive on medium containing 1 mM IPTG, yet were able to grow on similar medium lacking IPTG, were subjected to further growth sensitivity analysis

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as follows: Colonies from the plate lacking IPTG were manually picked and inoculated into individual wells of a 96 well culture dish containing LB + CB100 or Streptomycin 40, and were incubated for 16 hours at 30°C. These cultures were robotically diluted 1/100 into fresh medium and allowed to incubate for 4 hours at 37°C, after which they were subjected to serial dilutions in a 384 well array and then gridded onto media with and without 1 mM IPTG. After growth for 16 hours at 37°C, the arrays of serially diluted spots that resulted were compared between the two media. Clones that grew similarly at all dilutions on both media were scored as a negative and were no longer considered. Clones that grew on IPTG medium but failed to grow at the same serial dilution on the non-IPTG plate were given a score based on the differential, i.e. should the clone grow at a serial dilution of 10⁴ or less on the IPTG plate and grow at a serial dilution of 10⁸ or less on the IPTG plate, then the corresponding clone received a score of "4" representing the log difference in growth observed.

Following the identification of those vectors that, upon induction, negatively impacted *Pseudomonas aeruginosa* growth or proliferation, the inserts or nucleic acid fragments contained in those vectors were isolated for subsequent characterization. Vectors of interest were subjected to nucleic acid sequence determination.

Nucleic acids involved in proliferation of *E. faecalis* were identified as follows. Randomly generated fragments of genomic DNA were expressed from the vectors pEPEF3 or pEPEF14, which contain the CP25 or P59 promoter, respectively, regulated by the xyl operator/repressor. Should the genomic DNA downstream of the promoter contain, in an antisense orientation, at least a portion of a mRNA encoding a gene product involved in proliferation, then induction of expression from the promoter will result in detectable inhibition of proliferation.

A shotgun library of *E. faecalis* genomic fragments was cloned into the vector pEPEF3 or pEPEF14, which harbor xylose inducible promoters. The vector was linearized at a unique *SmaI* site immediately downstream of the promoter/operator. The linearized vector was treated with alkaline phosphatase to prevent reclosure of the linearized ends. Genomic DNA isolated from *E. faecalis* strain OG1RF was partially digested with DNase I and "blunt-ended" by incubating with T4 DNA polymerase. Random genomic fragments between 200 and 800 base pairs in length were selected by gel purification. The size-selected genomic fragments were added to the linearized and dephosphorylated vector at a molar ratio of 2 to 1, and ligated to form a shotgun library.

The ligated products were transformed into electrocompetent *E. coli* strain TOP10 cells (Invitrogen) and plated on LB medium with erythromycin (Erm) at 150 μ g/ml. Resulting colonies numbering 5 x 10⁵ or greater were scraped and combined, and were then subjected to plasmid purification.

The purified library was then transformed into electrocompetent *E. faecalis* strain OG1RF. Resulting transformants were plated on Todd-Hewitt (TH) agar with erythromycin at 10 µg/ml in

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order to generate 100 to 150 platings at 500 colonies per plating. The colonies were subjected to robotic picking and arrayed into wells of 384 well culture dishes. Each well contained 100 μ l of THB + Erm 10 μ g/ml. Inoculated 384 well dishes were incubated 16 hours at room temperature, and each well was robotically gridded onto solid TH agar + Erm with or without 5% xylose. Gridded plates were incubated 16 hours at 37°C, and then manually scored for arrayed colonies that were growth-compromised in the presence of xylose.

Arrayed colonies that were growth-sensitive on medium containing 5% xylose, yet were able to grow on similar medium lacking xylose, were subjected to further growth sensitivity analysis. Colonies from the plate lacking xylose were manually picked and inoculated into individual wells of a 96 well culture dish containing THB + Erm 10, and were incubated for 16 hours at 30°C. These cultures were robotically diluted 1/100 into fresh medium and allowed to incubate for 4 hours at 37°C, after which they were subjected to serial dilution on plates containing 5% xylose or plates lacking xylose. After growth for 16 hours at 37°C, the arrays of serially diluted spots that resulted were compared between the two media. Colonies that grew similarly on both media were scored as a negative and corresponding colonies were no longer considered. Colonies on xylose medium that failed to grow to the same serial dilution compared to those on the non-xylose plate were given a score based on the differential. For example, colonies on xylose medium that only grow to a serial dilution of -4 while they were able to grow to -8 on the non-xylose plate, then the corresponding transformant colony received a score of "4" representing the log difference in growth observed.

Following the identification of those vectors that, upon induction, negatively impacted *E. faecalis* growth or proliferation, the inserts or nucleic acid fragments contained in those expression vectors were isolated for subsequent characterization. The inserts in the vectors of interest were subjected to nucleotide sequence determination.

It will be appreciated that other restriction enzymes and other endonucleases or methodologies may be used to generate random genomic fragments. In addition, random genomic fragments may be generated by mechanical shearing. Sonication and nebulization are two such techniques commonly used for mechanical shearing of DNA.

EXAMPLE 2

Nucleotide Sequence Determination of Identified Clones Transribing Nucleic Acid Fragments with Detrimental Effects on Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae,

Pseudomonas aeruginosa or Enterococcus faecalis Proliferation

Plasmids from clones that received a dilution plating score of "2" or greater were isolated to obtain the genomic DNA insert responsible for growth inhibition as follows. *Staphylococcus aureus* were grown in standard laboratory media (LB or TB with 15 ug/ml Chloramphenicol to select for the plasmid). Growth was carried out at 37°C overnight in culture tubes or 2 ml deep well microtiter plates.

Lysis of Staphylococcus aureus was performed as follows. Cultures (2-5 ml) were centrifuged and the cell pellets resuspended in 1.5 mg/ml solution of lysostaphin (20 μ l/ml of original culture) followed by addition of 250 μ l of resuspension buffer (Qiagen). Alternatively, cell pellets were resuspended directly in 250 μ l of resuspension buffer (Qiagen) to which 5-20 μ l of a 1 mg/ml lysostaphin solution were added.

DNA was isolated using Qiagen miniprep kits or Wizard (Qiagen) miniprep kits according to the instructions provided by the manufacturer.

The genomic DNA inserts were amplified from the purified plasmids by PCR as follows.

 $1~\mu l$ of Qiagen purified plasmid was put into a total reaction volume of 25 $~\mu l$ Qiagen Hot Start PCR mix. For *Staphylococcus aureus*, the following primers were used in the PCR reaction:

20 pXyIT5F: CAGCAGTCTGAGTTATAAAATAG (SEQ ID NO: 1)

LexL TGTTTTATCAGACCGCTT (SEQ ID NO: 2)

Similar methods were conducted for Salmonella typhimurium and Klebsiella pneumoniae.

For Salmonella typhimurium and Klebsiella pneumoniae the following primers were used:

5' - TGTTTTATCAGACCGCTT- 3' (SEQ ID NO: 2) and

25 5'-ACAATTTCACACAGCCTC-3' (SEQ ID NO: 4)

PCR was carried out in a PE GenAmp with the following cycle times:

Step 1. 95° C 15 min

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Step 2. 94° C 45 sec

Step 3. 54° C 45 sec

30 Step 4, 72° C 1 minute

Step 5. Return to step 2, 29 times

Step 6. 72° C 10 minutes

Step 7. 4° C hold

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The PCR products were cleaned using Qiagen Qiaquick PCR plates according to the manufacturer's instructions.

For *Pseudomonas aeruginosa*, plasmids from transformant colonies that received a dilution plating score of "2" or greater were isolated to obtain the genomic DNA insert responsible for growth inhibition as follows. *Pseudomonas aeruginosa* were grown in standard laboratory media (LB with carbenicillin at 100 g/ml or Streptomycin 40 g/ml to select for the plasmid). Growth was carried out at 30°C overnight in 100 ul culture wells in microtiter plates. To amplify insert DNA 2 ul of culture were placed into 25 ul Qiagen Hot Start PCR mix. PCR reactions were in 96 well microtiter plates. For plasmid pEP5S the following primers were used in the PCR reaction:

T7L1+: GTCGGCGATATAGGCGCCAGCAACCG (SEO ID NO: 5)

pStrA3: ATAATCGAGCATGAGTATCATACG (SEQ ID NO: 6)

10 PCR was carried out in a PE GenAmp with the following cycle times:

Step 1. 95° C 15 min

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Step 2. 94° C 45 sec

Step 3. 54° C 45 sec

Step 4. 72° C 1 minute

15 Step 5. Return to step 2, 29 times

Step 6. 72° C 10 minutes

Step 7. 4° C hold

The PCR products were cleaned using Qiagen Qiaquick PCR plates according to the manufacturer's instructions.

20 The purified PCR products were then directly cycle sequenced with Qiagen Hot Start PCR mix. The following primers were used in the sequencing reaction:

T7/L2: ATGCGTCCGGCGTAGAGGAT (SEQ ID NO: 7)

PCR was carried out in a PE GenAmp with the following cycle times:

Step 1. 94° C 15 min

25 Step 2. 96° C 10 sec

Step 3. 50° C 5 sec

Step 4. 60 C 4 min

Step 5. Return to step 2, 24 times

Step 6. 4° C hold

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30 The PCR products were cleaned using Qiagen Qiaquick PCR plates according to the manufacturer's instructions.

For *E. faecalis*, plasmids from transformant colonies that received a dilution plating score of "2" or greater were isolated to obtain the genomic DNA insert responsible for growth inhibition as follows. *E. faecalis* were grown in THB 10 μ g/ml Erm at 30°C overnight in 100 μ g culture wells in microtiter plates. To amplify insert DNA 2 μ g of culture were placed into 25 μ g Qiagen Hot Start

PCR mix. PCR reactions were in 96 well microtiter plates. The following primers were used in the PCR reaction:

pXyIT5: CAGCAGTCTGAGTTATAAAATAG (SEQ ID NO: 1) and the pEP/pAK1 primer.

5 PCR was carried out in a PE GenAmp with the following cycle times:

Step 1. 95° C 15 min

Step 2. 94° C 45 sec

Step 3. 54° C 45 sec

Step 4. 72° C 1 minute

10 Step 5. Return to step 2, 29 times

Step 6. 72° C 10 minutes

Step 7. 4° C hold

The PCR products were cleaned using Qiagen Qiaquick PCR plates according to the manufacturer's instructions.

15 The purified PCR products were then directly cycle sequenced with Qiagen Hot Start PCR mix. The following primers were used in the PCR reaction:

pXylT5: CAGCAGTCTGAGTTATAAAATAG (SEQ ID NO: 1)

PCR was carried out in a PE GenAmp with the following cycle times:

Step 1. 94° C 15 min

20 Step 2. 96° C 10 sec

Step 3. 50° C 5 sec

Step 4, 60° C 4 min

Step 5. Return to step 2, 24 times

Step 6. 4° C hold

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25 The PCR products were cleaned using Qiagen Qiaquick PCR plates according to the manufacturer's instructions.

The amplified genomic DNA inserts from each of the above procedures were subjected to automated sequencing. Sequence identification numbers (SEQ ID NOs) and clone names for the identified inserts are listed in Table IA and discussed below.

EXAMPLE 3

Comparison Of Isolated Nucleic Acids to Known Sequences

The nucleotide sequences of the subcloned fragments from Staphylococcus aureus,

Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa or Enterococcus

faecalis obtained from the expression vectors discussed above were compared to known sequences
from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas
aeruginosa or Enterococcus faecalis and other microorganisms as follows. First, to confirm that

each clone originated from one location on the chromosome and was not chimeric, the nucleotide sequences of the selected clones were compared against the Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa or Enterococcus faecalis genomic sequences to align the clone to the correct position on the chromosome. The NCBI BLASTN v 2.0.9 program was used for this comparison, and the incomplete Staphylococcus aureus genomic sequences licensed from TIGR, as well as the NCBI nonredundant GenBank database were used as the source of genomic data. Salmonella typhimurium sequences were compared to sequences available from the Genome Sequencing Center (http://genome.wustl.edu/gsc/salmonella.shtml),and the Sanger Centre (http://www.sanger.ac.uk/projects/S__typhi). Pseudomonas aeruginosa sequences were compared to a proprietary database and the NCBI GenBank database. The E. faecalis sequences were compared to a proprietary database.

The BLASTN analysis was performed using the default parameters except that the filtering was turned off. No further analysis was performed on inserts which resulted from the ligation of multiple fragments.

In general, antisense molecules and their complementary genes are identified as follows. First, all possible full length open reading frames (ORFs) are extracted from available genomic databases. Such databases include the GenBank nonredundant (nr) database, the unfinished genome database available from TIGR and the PathoSeq database developed by Incyte Genomics. The latter database comprises over 40 annotated bacterial genomes including complete ORF analysis. If databases are incomplete with regard to the bacterial genome of interest, it is not necessary to extract all ORFs in the genome but only to extract the ORFs within the portions of the available genomic sequences which are complementary to the clones of interest. Computer algorithms for identifying ORFs, such as GeneMark, are available and well known to those in the art. Comparison of the clone DNA to the complementary ORF(s) allows determination of whether the clone is a sense or antisense clone. Furthermore, each ORF extracted from the database can be compared to sequences in well annotated databases including the GenBank (nr) protein database, SWISSPROT and the like. A description of the gene or of a closely related gene in a closely related microorganism is often available in these databases. Similar methods are used to identify antisense clones corresponding to genes encoding non-translated RNAs.

In order to generate the gene identification data compiled in Table IB, each of the cloned nucleic acid sequences discussed above corresponding to SEQ ID NO.s 8-3795 was used to identify the corresponding Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa or Enterococcus faecalis ORFs in the PathoSeq v.4.1 (March 2000 release) database of microbial genomic sequences. For this purpose, the NCBI BLASTN 2.0.9 computer algorithm was used. The default parameters were used except that filtering was turned off. The default parameters for the BLASTN and BLASTX analyses were:

Expectation value (e)=10

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Alignment view options: pairwise

Filter query sequence (DUST with BLASTN, SEG with others)=T

Cost to open a gap (zero invokes behavior)=0

Cost to extend a gap (zero invokes behavior)=0

5 X dropoff value for gapped alignment (in bits) (zero invokes behavior)=0

Show GI's in deflines=F

Penalty for a nucleotide mismatch (BLASTN only)=-3

Reward for a nucleotide match (BLASTN only)=1

Number of one-line descriptions (V)=500

Number of alignments to show (B)=250

Threshold for extending hits=default

Perform gapped alignment (not available with BLASTX)=T

Query Genetic code to use=1

DB Genetic code (for TBLASTInx) only=1

Number of processors to use=1

SeaAlign file

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Believe the query defline=F

Matrix=BLOSUM62

Word Size= default

Effective length of the database (use zero for the real size)=0

Number of best hits from a region to keep=100

Length of region used to judge hits=20

Effective length of the search space (use zero for the real size)=0

Query strands to search against database (for BLAST[nx] and TBLASTX), 3 is both, 1 is top, 2 is bottom=3

Produce HTML output=F

Alternatively, ORFs were identified and refined by conducting a survey of the public and private data sources. Full-length gene protein and nucleotide sequences for these organisms were assembled from various sources. For *Pseudomonas aeruginosa*, gene sequences were adopted from the Pseudomonas genome sequencing project (downloaded from http://www.pseudomonas.com). For *Klebsiella pneumoniae*, *Staphylococcus aureus*, *Streptococcus pneumoniae* and *Salmonella*

typhi, genomic sequences from PathoSeq v 4.1 (Mar 2000 release) was reanalyzed for ORFs using the gene finding software GeneMark v 2.4a, which was purchased from GenePro Inc. 451 Bishop St., N.W., Suite B, Atlanta, GA, 30318, USA.

Antisense clones were identified as those clones for which transcription from the inducible promoter would result in the expression of an RNA antisense to a complementary ORF, intergenic or intragenic sequence. Those clones containing single inserts and that caused growth sensitivity upon induction are listed in Table IA. ORFs complementary to the antisense nucleic acids, and their encoded polypeptides, are listed in Table IB.

The gene descriptions in the PathoSeq database derive from annotations available in the public sequence databases described above. Where a clone was found to share significant sequence identity to two or more adjacent ORFs, it was listed once for each ORF and the PathoSeq information for each ORF was compiled in Table IB.

Table IA lists the SEQ ID NOs. and clone names of the inserts which inhibited proliferation and the organism in which the clone was identified. This information was used to identify the

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ORFs (SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012) whose gene products (SEQ ID NOs. 3801-3805, 4861-5915, 10013-14110) were inhibited by the nucleic acids comprising the nucleotide sequences of SEQ ID NOs. 8-3795. Table IB lists the clone name, the SEQ ID NO. of the antisense clone (in the column labelled Clone SEQ ID), the PathoSeq Locus containing the clone, the SEQ ID of the ORF identified in PathoSeq (in the column labelled Gene Seq ID (protein), the refined full length gene (column labelled genemarked gene), and the SEQ ID NO of the protein encoded by the refined full length gene (column labelled full length ORF protein SEQ ID).

Table IC provides a cross reference between PathoSeq Gene Locus listed in Table IB, the SEQ ID NOs. of the PathoSeq proteins and the SEQ ID NOs. of the nucleic acids which encode them.

It will be appreciated that ORFs may also be identified using databases other than PathoSeq. For example, the ORFs may be identified using the methods described in U.S. Provisional Patent Application Serial Number 60/191,078, filed March 21, 2000.

EXAMPLE 4

Identification of Genes and their Corresponding Operons Affected by Antisense Inhibition

Once the genes involved in Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa or Enterococcus faecalis proliferation are identified as described above, the operons in which these genes lie may be identified by comparison with known microbial genomes. Since bacterial genes are transcribed in a polycistronic manner, the antisense inhibition of a single gene in an operon might affect the expression of all the other genes on the operon or the genes downstream from the single gene identified. Accordingly, each of the genes contained within an operon may be analyzed for their effect on proliferation.

Operons are predicted by looking for all adjacent genes in a genomic region that lie in the same orientation with no large noncoding gaps in between. First, full-length ORFs complementary to the antisense molecules are identified as described above. Adjacent ORFs are then identified and their relative orientation determined either by directly analyzing the genomic sequences surrounding the ORFs complementary to the antisense clones or by extracting adjacent ORFs from the collection obtained through whole genome ORF analysis described above followed by ORF alignment. Operons predicted in this way may be confirmed by comparison to the arrangement of the homologous nucleic acids in the *Bacillus subtilis* complete genome sequence, as reported by the genome database compiled at Institut Pasteur Subtilist Release R15.1 (June 24, 1999) which can be found at http://bioweb.pasteur.fr/GenoList/SubtiList/. The *Bacillus subtilis* genome is the only fully sequenced and annotated genome from a Gram-positive microorganism, and appears to have a high level of similarity to *Staphylococcus aureus* both at the level of conservation of gene sequence and genomic organization including operon structure. Operons for *Salmonella typhimurium* and *Klebsiella pneumoniae* may be identified by comparison with *E. coli, Haemophilus*, or

Pseudomonas sequences. The *Pseudomonas aeruginosa* web site (http://www.pseudomonas.com) can also be used to help predict operon organization in this bacterium.

Extensive DNA sequences of Salmonella typhimurium are available through the Salmonella Genome Center (Washington University, St. Louis, MO) the Sanger Centre (United Kingdom) and the PathoSeq database (Incyte). Annotation of some of the DNA sequences in some of the aforementioned databases is lacking, but comparisons may be made to E. coli using tools such as BLASTX.

Public or proprietary databases may be used to analyzed *E. faecalis* sequences as well as sequences from the organisms listed above.

The results of such an analysis as applied to clone number S1M10000001A05 from Staphylococcus aureus are listed in Table II. Table II lists the SEQ ID NOs. of the Staphylococcus aureus genes involved in proliferation, the SEQ ID NOs. of the proteins encoded by these genes, and the clone name containing the nucleic acid which inhibits Staphylococcus aureus proliferation. In addition, Table II lists those other genes located on the operon included in the Staphylococcus aureus genomic sequence determined as described above. For each of the genes described in Table II, the microorganism containing the most closely related homolog, identified in one of the public databases, is also indicated in Table II.

TABLE II

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DNA Seq ID	Protein Seq ID	Molecule number	Clone пате	Gene	Organism used for identification of gene
3796	3801	SaXA001	S1M10000001A05	ytmI	B. subtilis
3797	3802			nirR	S. carnosus
3798	3803			nirB	S. carnosus
3799	3804			nirD	S. carnosus
3800	3805			sirB	S. carnosus

The preceding analyses may be conducted for each of the sequences which are listed in Table IA which inhibit proliferation and the ORFs listed in Table IB and Table IC. Once the full length ORFs and/or the operons containing them have been identified using the methods described above, they can be obtained from a genomic library by performing a PCR amplification using primers at each end of the desired sequence. Those skilled in the art will appreciate that a comparison of the ORFs to homologous sequences in other cells or microorganisms will facilitate confirmation of the start and stop codons at the ends of the ORFs.

In some embodiments, the primers may contain restriction sites which facilitate the insertion of the gene or operon into a desired vector. For example, the gene may be inserted into an expression vector and used to produce the proliferation-required protein as described below. Other methods for obtaining the full length ORFs and/or operons are familiar to those skilled in the art.

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For exmaple, natural restriction sites may be employed to insert the full length ORFs and/or operons into a desired vector.

EXAMPLE 5

Identification of Individual Genes within an Operon Required for Proliferation

The following example illustrates a method for determining if a targeted gene within an operon is required for cell proliferation by replacing the targeted allele in the chromosome with an in-frame deletion of the coding region of the targeted gene.

Deletion inactivation of a chromosomal copy of a gene in Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi can be accomplished by integrative gene replacement. The principles of this method were described in Xia, M., et al. 1999 Plasmid 42:144-149 and Hamilton, C. M., et al. 1989. J. Bacteriol. 171: 4617-4622. A similar gene disruption method is available for Pseudomonas aeruginosa, except the counter selectable marker is sacB (Schweizer, H. P., Klassen, T. and Hoang, T. (1996) Mol. Biol. of Pseudomonas. ASM press, 229-237). In this approach, a mutant allele of the targeted gene is constructed by way of an in-frame deletion and introduced into the chromosome using a suicide vector. This results in a tandem duplication comprising a deleted (null) allele and a wild type allele of the target gene. Cells in which the vector sequences have been deleted are isolated using a counter-selection technique. Removal of the vector sequence from the chromosomal insertion results in either restoration of the wild-type target sequence or replacement of the wild type sequence with the deletion (null) allele. E. faecalis genes can be disrupted using a suicide vector that contains an internal fragment to a gene of interest. With the appropriate selection this plasmid will homologously recombine into the chromosome (Nallapareddy, S. R., X. Qin, G. M. Weinstock, M. Hook, B. E. Murray. 2000. Infect. Immun. 68:5218-5224).

The resultant population of Staphylococcus aureus; Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi colonies can then be evaluated to determine whether the target sequence is required for proliferation by PCR amplification of the affected target sequence. If the targeted gene is not required for proliferation, then PCR analysis will show that roughly equal numbers of colonies have retained either the wild-type or the mutant allele. If the targeted gene is required for proliferation, then only wild-type alleles will be recovered in the PCR analysis.

The method of cross-over PCR is used to generate the mutant allele by amplification of nucleotide sequences flanking but not including the coding region of the gene of interest, using specifically designed primers such that overlap between the resulting two PCR amplification products allows them to hybridize. Further PCR amplification of this hybridization product using

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primers representing the extreme 5' and 3' ends can produce an amplification product containing an in-frame deletion of the coding region but retaining substantial flanking sequences.

For Staphylococcus aureus, this amplification product is subcloned into the suicide vector pSA3182 (Xia, M., et al. 1999 Plasmid 42:144-149) which is host-dependent for autonomous replication. This vector includes a tetC tetracycline-resistance marker and the origin of replication of the well-known Staphylococcus aureus plasmid pT181 (Mojumdar, M and Kahn, S.A., Characterisation of the Tetracycline Resistance Gene of Plasmid pT181, J. Bacteriol. 170: 5522 (1988)). The vector lacks the repC gene which is required for autonomous replication of the vector at the pT181 origin. This vector can be propagated in a Staphylococcus aureus host strain such as SA3528, which expresses repC in trans. Once the amplified truncated target gene sequence is cloned and propagated in the pSA3182 vector, it can then be introduced into a repC minus strain such as RN4220 (Kreiswirth, B.N. et al., The Toxic Shock Syndrome Exotoxin Structural Gene is Not Detectably Transmitted by a Prophage, Nature 305:709-712 (1983)) by electroporation with selection for tetracycline resistance. In this strain, the vector must integrate by homologous recombination at the targeted gene in the chromosome to impart drug resistance. This results in a inserted truncated copy of the allele, followed by pSA3182 vector sequence, and finally an intact and functional allele of the targeted gene.

Once a tetracycline resistant Staphylococcus aureus strain is isolated using the above technique and shown to include truncated and wild-type alleles of the targeted gene as described above, a second plasmid, pSA7592 (Xia, M., et al. 1999 Plasmid 42:144-149) is introduced into the strain by electroporation. This gene includes an erythromycin resistance gene and a repC gene that is expressed at high levels. Expression of repC in these transformants is toxic due to interference of normal chromosomal replication at the integrated pT181 origin of replication. This selects for strains that have removed the vector sequence by homologous recombination, resulting in either of two outcomes: The selected cells either possess a wild-type allele of the targeted gene or a gene in which the wild-type allele has been replaced by the engineered in-frame deletion of the truncated allele.

PCR amplification can be used to determine the genetic outcome of the above process in the resulting erythromycin resistant, tet sensitive transformant colonies. If the targeted gene is not required for cellular replication, then PCR evidence for both wild-type and mutant alleles will be found among the population of resultant transformants. However, if the targeted gene is required for cellular proliferation, then only the wild-type form of the gene will be evident among the resulting transformants.

Similarly, for Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa or Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi the PCR products containing the mutant allele of the

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target sequence may be introduced into an appropriate knockout vector and cells in which the wild type target has been disrupted are selected using the appropriate methodology.

The above methods have the advantage that insertion of an in-frame deletion mutation is far less likely to cause downstream polar effects on genes in the same operon as the targeted gene. However, it will be appreciated that other methods for disrupting Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi genes which are familiar to those skilled in the art may also be used.

Each gene in the operon may be disrupted using the methodology above to determine whether it is required for proliferation.

EXAMPLE 6

Expression of the Proteins Encoded by Genes Identified as

Required for Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae,

Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis,

Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi Proliferation

The following is provided as one exemplary method to express the proliferation-required proteins idenfied as described above. The proliferation-required proteins may be expressed using any of the bacterial, insect, yeast, or mammalian expression systems known in the art. In some embodiments, the proliferation-required proteins encoded by the identified nucleotide sequences described above (including the proteins of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110 encoded by the nucleic acids of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012 are expressed using expression systems designed either for E. coli or for Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi, First, the initiation and termination codons for the gene are identified. If desired, methods for improving translation or expression of the protein are well known in the art. For example, if the nucleic acid encoding the polypeptide to be expressed lacks a methionine codon to serve as the initiation site, a strong Shine-Delgarno sequence, or a stop codon, these nucleotide sequences can be added. Similarly, if the identified nucleic acid lacks a transcription termination signal, this nucleotide sequence can be added to the construct by, for example, splicing out such a sequence from an appropriate donor sequence. In addition, the coding sequence may be operably linked to a strong constitutive promoter or an inducible promoter if desired. The identified nucleic acid or portion thereof encoding the polypeptide to be expressed is obtained by, for example, PCR from the bacterial expression vector or genome using oligonucleotide primers complementary to the identified nucleic acid or portion thereof and containing restriction endonuclease sequences appropriate for inserting the coding sequences into the vector such that the coding sequences can be expressed from the vector's promoter. Alternatively, other conventional cloning techniques may be used to place the coding sequence under the control of

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the promoter. In some embodiments, a termination signal may be located downstream of the coding sequence such that transcription of the coding sequence ends at an appropriate position.

Several expression vector systems for protein expression in *E. coli* are well known and available to those knowledgeable in the art. The coding sequence may be inserted into any of these vectors and placed under the control of the promoter. The expression vector may then be transformed into DH5α or some other *E. coli* strain suitable for the over expression of proteins.

Alternatively, an expression vector encoding a protein required for proliferation of Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa. Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae. Helicobacter pylori, or Salmonella typhi may be introduced into Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli. Enterococcus faecalis. Haemophilus influenzae. Helicobacter pylori. ox Salmonella typhi . Protocols for introducing nucleic acids into these organisms are well known in the art. For example, the protocols described in J.C.Lee "Electroporation of Staphylococci" from Methods in Molecular Biology vol 47: Electroporation Protocols for Microorganisms Edited by: J.A. Nickoloff Humana Press Inc., Totowa, NJ. pp209-216, may be used to introduce nucleic acids into Staphylococcus aureus. Nucleic acids may also be introduced into Salmonella typhimurium, Klebsiella pneumoniae. Pseudomonas aeruginosa or Enterococcus faecalis using methods familiar to those skilled in the art. Positive transformants are selected after growing the transformed cells on plates containing an antibiotic to which the vector confers resistance. In one embodiment, Staphylococcus aureus is transformed with an expression vector in which the coding sequence is operably linked to the T5 promoter containing a xylose operator such that expression of the encoded protein is inducible with xvlose.

In one embodiment, the protein is expressed and maintained in the cytoplasm as the native sequence. In an alternate embodiment, the expressed protein can be modified to include a protein tag that allows for differential cellular targeting, such as to the periplasmic space of Gram-negative or Gram-positive expression hosts or to the exterior of the cell (i.e., into the culture medium). In some embodiments, the osmotic shock cell lysis method described in Chapter 16 of Current Protocols in Molecular Biology, Vol. 2, (Ausubel, et al., Eds.) John Wiley & Sons, Inc. (1997) may be used to liberate the polypeptide from the cell. In still another embodiment, such a protein tag could also facilitate purification of the protein from either fractionated cells or from the culture medium by affinity chromatography. Each of these procedures can be used to express a proliferation-required protein.

Expressed proteins, whether in the culture medium or liberated from the periplasmic space or the cytoplasm, are then purified or enriched from the supernatant using conventional techniques such as ammonium sulfate precipitation, standard chromatography, immunoprecipitation, immunochromatography, size exclusion chromatography, ion exchange chromatography, and HPLC.

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Alternatively, the polypeptide may be secreted from the host cell in a sufficiently enriched or pure state in the supernatant or growth media of the host cell to permit it to be used for its intended purpose without further enrichment. The purity of the protein product obtained can be assessed using techniques such as SDS PAGE, which is a protein resolving technique well known to those skilled in the art. Coomassie, silver staining or staining with an antibody are typical methods used to visualize the protein of interest.

Antibodies capable of specifically recognizing the protein of interest can be generated using synthetic peptides using methods well known in the art. See, Antibodies: A Laboratory Manual, (Harlow and Lane, Eds.) Cold Spring Harbor Laboratory (1988). For example, 15-mer peptides having an amino acid sequence encoded by the appropriate identified gene sequence of interest or portion thereof can be chemically synthesized. The synthetic peptides are injected into mice to generate antibodies to the polypeptide encoded by the identified nucleic acid sequence of interest or portion thereof. Alternatively, samples of the protein expressed from the expression vectors discussed above can be purified and subjected to amino acid sequencing analysis to confirm the identity of the recombinantly expressed protein and subsequently used to raise antibodies. An Example describing in detail the generation of monoclonal and polyclonal antibodies appears in Example 7.

The protein encoded by the identified nucleic acid of interest or portion thereof can be purified using standard immunochromatography techniques. In such procedures, a solution containing the secreted protein, such as the culture medium or a cell extract, is applied to a column having antibodies against the secreted protein attached to the chromatography matrix. The secreted protein is allowed to bind the immunochromatography column. Thereafter, the column is washed to remove non-specifically bound proteins. The specifically-bound secreted protein is then released from the column and recovered using standard techniques. These procedures are well known in the art.

In an alternative protein purification scheme, the identified nucleic acid of interest or portion thereof can be incorporated into expression vectors designed for use in purification schemes employing chimeric polypeptides. In such strategies the coding sequence of the identified nucleic acid of interest or portion thereof is inserted in-frame with the gene encoding the other half of the chimera. The other half of the chimera can be maltose binding protein (MBP) or a nickel binding polypeptide encoding sequence. A chromatography matrix having maltose or nickel attached thereto is then used to purify the chimeric protein. Protease cleavage sites can be engineered between the MBP gene or the nickel binding polypeptide and the identified expected gene of interest, or portion thereof. Thus, the two polypeptides of the chimera can be separated from one another by protease digestion.

One useful expression vector for generating maltose binding protein fusion proteins is pMAL (New England Biolabs), which encodes the *malE* gene. In the pMal protein fusion system, the cloned gene is inserted into a pMal vector downstream from the *malE* gene. This results in the expression of an MBP-fusion protein. The fusion protein is purified by affinity chromatography. These techniques as described are well known to those skilled in the art of molecular biology.

EXAMPLE 7

Production of an Antibody to an isolated Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa "Enterococcus faecalis, Escherichia coli, Enterococcus

faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi Protein

Substantially pure protein or polypeptide (including one of the polypeptides of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110) is isolated from the transformed cells as described in Example 6. The concentration of protein in the final preparation is adjusted, for example, by concentration on a 10,000 molecular weight cut off AMICON filter device (Millipore, Bedford, MA), to the level of a few micrograms/ml. Monoclonal or polyclonal antibody to the protein can then be prepared as follows:

Monoclonal Antibody Production by Hybridoma Fusion

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Monoclonal antibody to epitopes of any of the peptides identified and isolated as described can be prepared from murine hybridomas according to the classical method of Kohler, G. and Milstein, C., Nature 256:495 (1975) or any of the well-known derivative methods thereof. Briefly, a mouse is repetitively inoculated with a few micrograms of the selected protein or peptides derived therefrom over a period of a few weeks. The mouse is then sacrificed, and the antibody-producing cells of the spleen isolated. The spleen cells are fused by means of polyethylene glycol with mouse myeloma cells, and the excess unfused cells are destroyed by growth of the system on selective medium comprising aminopterin (HAT medium). The successfully-fused cells are diluted and aliquots of the dilution placed in wells of a microtiter plate where growth of the culture is continued. Antibody-producing clones are identified by detection of antibody in the supernatant fluid of the wells by immunoassay procedures, such as ELISA, as described by Engvall, E., "Enzyme immunoassay ELISA and EMIT," Meth. Enzymol. 70:419 (1980), and derivative methods thereof. Selected positive clones can be expanded and their monoclonal antibody product harvested for use. Detailed procedures for monoclonal antibody production are described in Davis, L. et al. Basic Methods in Molecular Biology Elsevier, New York. Section 21-2.

Polyclonal Antibody Production by Immunization

Polyclonal antiserum containing antibodies to heterogeneous epitopes of a single protein or a peptide can be prepared by immunizing suitable animals with the expressed protein or peptides derived therefrom described above, which can be unmodified or modified to enhance immunogenicity. Effective polyclonal antibody production is affected by many factors related both to the antigen and the host species. For example, small molecules tend to be less immunogenic than larger molecules and can require the use of carriers and adjuvant. Also, host animals vary in response to site of inoculations and dose, with both inadequate or excessive doses of antigen resulting in low titer antisera. Small doses (ng level) of antigen administered at multiple intradermal sites appears to be most reliable. An effective immunization protocol for rabbits can be found in Vaitukaitis, J. et al. J. Clin. Endocrinol. Metab. 33:988-991 (1971).

Booster injections can be given at regular intervals, and antiserum harvested when antibody titer thereof, as determined semi-quantitatively, for example, by double immunodiffusion in agar against known concentrations of the antigen, begins to fall. See, for example, Ouchterlony, O. et al., Chap. 19 in: **Handbook of Experimental Immunology** D. Wier (ed) Blackwell (1973). Plateau concentration of antibody is usually in the range of 0.1 to 0.2 mg/ml of serum (about 12 μM). Affinity of the antisera for the antigen is determined by preparing competitive binding curves, as described, for example, by Fisher, D., Chap. 42 in: **Manual of Clinical Immunology**, 2d Ed. (Rose and Friedman, Eds.) Amer. Soc. For Microbiol., Washington, D.C. (1980).

Antibody preparations prepared according to either protocol are useful in quantitative immunoassays which determine concentrations of antigen-bearing substances in biological samples; they are also used semi-quantitatively or qualitatively to identify the presence of antigen in a biological sample. The antibodies can also be used in therapeutic compositions for killing bacterial cells expressing the protein.

EXAMPLE 8

Screening Chemical Libraries

A. Protein-Based Assays

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Having isolated and expressed bacterial proteins shown to be required for bacterial proliferation, the present invention further contemplates the use of these expressed target proteins in assays to screen libraries of compounds for potential drug candidates. The generation of chemical libraries is well known in the art. For example, combinatorial chemistry can be used to generate a library of compounds to be screened in the assays described herein. A combinatorial chemical library is a collection of diverse chemical compounds generated by either chemical synthesis or biological synthesis by combining a number of chemical "building block" reagents. For example, a linear combinatorial chemical library such as a polypeptide library is formed by combining amino acids in every possible combination to yield peptides of a given length. Millions of chemical compounds theoretically can be synthesized through such combinatorial mixings of chemical building blocks. For example, one commentator observed that the systematic, combinatorial mixing of 100 interchangeable chemical building blocks results in the theoretical synthesis of 100 million tetrameric compounds or 10 billion pentameric compounds. (Gallop et al., "Applications of Combinatorial Technologies to Drug Discovery, Background and Peptide Combinatorial Libraries," Journal of Medicinal Chemistry, Vol. 37, No. 9, 1233-1250 (1994). Other chemical libraries known to those in the art may also be used, including natural product libraries.

Once generated, combinatorial libraries can be screened for compounds that possess desirable biological properties. For example, compounds which may be useful as drugs or to develop drugs would likely have the ability to bind to the target protein identified, expressed and purified as discussed above. Further, if the identified target protein is an enzyme, candidate compounds would likely interfere with the enzymatic properties of the target protein. For example, the enzymatic function of a

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target protein may be to serve as a protease, nuclease, phosphatase, dehydrogenase, transporter protein, transcriptional enzyme, and any other type of enzyme known or unknown. Thus, the present invention contemplates using the protein products described above to screen combinatorial chemical libraries.

In one example, the target protein is a scrine protease and the substrate of the enzyme is known. The present example is directed towards the analysis of libraries of compounds to identify compounds that function as inhibitors of the target enzyme. First, a library of small molecules is generated using methods of combinatorial library formation well known in the art. U.S. Patent Nos. 5,463,564 and 5,574, 656, to Agrafiotis, et al., entitled "System and Method of Automatically Generating Chemical Compounds with Desired Properties," are two such teachings. Then the library compounds are screened to identify those compounds that possess desired structural and functional properties. U.S. Patent No. 5,684,711, also discusses a method for screening libraries.

To illustrate the screening process, the target polypeptide and chemical compounds of the library are combined with one another and permitted to interact with one another. A labeled substrate is added to the incubation. The label on the substrate is such that a detectable signal is emitted from the products of the substrate molecules that result from the activity of the target polypeptide. The emission of this signal permits one to measure the effect of the combinatorial library compounds on the enzymatic activity of target enzymes by comparing it to the signal emitted in the absence of combinatorial library compounds. The characteristics of each library compound are encoded so that compounds demonstrating activity against the enzyme can be analyzed and features common to the various compounds identified can be isolated and combined into future iterations of libraries.

Once a library of compounds is screened, subsequent libraries are generated using those chemical building blocks that possess the features shown in the first round of screen to have activity against the target enzyme. Using this method, subsequent iterations of candidate compounds will possess more and more of those structural and functional features required to inhibit the function of the target enzyme, until a group of enzyme inhibitors with high specificity for the enzyme can be found. These compounds can then be further tested for their safety and efficacy as antibiotics for use in mammals.

It will be readily appreciated that this particular screening methodology is exemplary only. Other methods are well known to those skilled in the art. For example, a wide variety of screening techniques are known for a large number of naturally-occurring targets when the biochemical function of the target protein is known. For example, some techniques involve the generation and use of small peptides to probe and analyze target proteins both biochemically and genetically in order to identify and develop drug leads. Such techniques include the methods described in PCT publications No. WO9935494, WO9819162, WO9954728. Other techniques utilize natural product libraries or libraries of larger molecules such as proteins.

It will be appreciated that the above protein-based assays may be performed with any of the proliferation-required polypeptides from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi (including the polypeptides of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110) or portions thereof. In addition, the above protein-based assays may be performed with homologous polypeptides or portions thereof.

B. Cell-Based Assays

Current cell-based assays used to identify or to characterize compounds for drug discovery and development frequently depend on detecting the ability of a test compound to modulate the activity of a target molecule located within a cell or located on the surface of a cell. An advantage of cell-based assays is that they allow the effect of a compound on a target molecule's activity to be detected within the physiologically relevant environment of the cell as opposed to an in vitro environment. Most often such target molecules are proteins such as enzymes, receptors and the like. However, target molecules may also include other molecules such as DNAs, lipids, carbohydrates and RNAs including messenger RNAs, ribosomal RNAs, tRNAs, regulatory RNAs and the like. A number of highly sensitive cell-based assay methods are available to those of skill in the art to detect binding and interaction of test compounds with specific target molecules. However, these methods are generally not highly effective when the test compound binds to or otherwise interacts with its target molecule with moderate or low affinity. In addition, the target molecule may not be readily accessible to a test compound in solution, such as when the target molecule is located inside the cell or within a cellular compartment. Thus, current cell-based assay methods are limited in that they are not effective in identifying or characterizing compounds that interact with their targets with moderate to low affinity or compounds that interact with targets that are not readily accessible.

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The cell-based assays methods of the present invention have substantial advantages over current cell-based assays. These advantages derive from the use of sensitized cells in which the level or activity of at least one proliferation-required gene product (the target molecule) has been specifically reduced to the point where the presence or absence of its function becomes a rate-determining step for cellular proliferation. Bacterial, fungal, plant, or animal cells can all be used with the present method. Such sensitized cells become much more sensitive to compounds that are active against the affected target molecule. Thus, cell-based assays of the present invention are capable of detecting compounds exhibiting low or moderate potency against the target molecule of interest because such compounds are substantially more potent on sensitized cells than on non-sensitized cells. The effect may be such that a test compound may be two to several times more potent, at least 10 times more potent, at least 20 times more potent, at least 50 times more potent, at least 100 times more potent, at least 1000 times more potent, or even more than 1000 times more potent when tested on the sensitized cells as compared to the non-sensitized cells. The

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proliferation-required nucleic acids or polypeptides from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi, or portions thereof, may be employed in any of the cell-based assays described herein. Similarly, homologous coding nucleic acids, homologous antisense nucleic acids, or homologous polypeptides or portions of the homologous nucleic acids or homologous polypeptides, may be employed in any of the cell-based assays described herein.

Due in part to the increased appearance of antibiotic resistance in pathogenic microorganisms and to the significant side-effects associated with some currently used antibiotics, novel antibiotics acting at new targets are highly sought after in the art. Yet, another limitation in the current art related to cell-based assays is the problem of repeatedly identifying hits against the same kinds of target molecules in the same limited set of biological pathways. This may occur when compounds acting at such new targets are discarded, ignored or fail to be detected because compounds acting at the "old" targets are encountered more frequently and are more potent than compounds acting at the new targets. As a result, the majority of antibiotics in use currently interact with a relatively small number of target molecules within an even more limited set of biological pathways.

The use of sensitized cells of the current invention provides a solution to the above problem in two ways. First, desired compounds acting at a target of interest, whether a new target or a previously known but poorly exploited target, can now be detected above the "noise" of compounds acting at the "old" targets due to the specific and substantial increase in potency of such desired compounds when tested on the sensitized cells of the current invention. Second, the methods used to sensitize cells to compounds acting at a target of interest may also sensitize these cells to compounds acting at other target molecules within the same biological pathway. For example, expression of an antisense molecule to a gene encoding a ribosomal protein is expected to sensitize the cell to compounds acting at that ribosomal protein and may also sensitize the cells to compounds acting at any of the ribosomal components (proteins or rRNA) or even to compounds acting at any target which is part of the protein synthesis pathway. Thus an important advantage of the present invention is the ability to reveal new targets and pathways that were previously not readily accessible to drug discovery methods.

Sensitized cells of the present invention are prepared by reducing the activity or level of a target molecule. The target molecule may be a gene product, such as an RNA or polypeptide produced from the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi (including a gene product produced from the nucleic acids of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, such as the polypeptides of SEO ID NOs.: 3801-3805, 4861-

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5915, 10013-14110) or from homologous nucleic acids. For example, the target molecule may be one of the polypeptides of SEQ ID NOs. 3801-3805, 4861-5915, 10013-14110 or a homologous polypeptide. Alternatively, the target may be a gene product such as an RNA or polypeptide which is produced from a sequence within the same operon as the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi or from homologous nucleic acids. In addition, the target may be an RNA or polypeptide in the same biological pathway as the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi or from homologous nucleic acids. Such biological pathways include, but are not limited to, enzymatic, biochemical and metabolic pathways as well as pathways involved in the production of cellular structures such the cell wall.

Current methods employed in the arts of medicinal and combinatorial chemistries are able to make use of structure-activity relationship information derived from testing compounds in various biological assays including direct binding assays and cell-based assays. Occasionally compounds are directly identified in such assays that are sufficiently potent to be developed as drugs. More often, initial hit compounds exhibit moderate or low potency. Once a hit compound is identified with low or moderate potency, directed libraries of compounds are synthesized and tested in order to identify more potent leads. Generally these directed libraries are combinatorial chemical libraries consisting of compounds with structures related to the hit compound but containing systematic variations including additions, subtractions and substitutions of various structural features. When tested for activity against the target molecule, structural features are identified that either alone or in combination with other features enhance or reduce activity. This information is used to design subsequent directed libraries containing compounds with enhanced activity against the target molecule. After one or several iterations of this process, compounds with substantially increased activity against the target molecule are identified and may be further developed as drugs. This process is facilitated by use of the sensitized cells of the present invention since compounds acting at the selected targets exhibit increased potency in such cell-based assays, thus; more compounds can now be characterized providing more useful information than would be obtained otherwise.

Thus, it is now possible using cell-based assays of the present invention to identify or characterize compounds that previously would not have been readily identified or characterized including compounds that act at targets that previously were not readily exploited using cell-based assays. The process of evolving potent drug leads from initial hit compounds is also substantially

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improved by the cell-based assays of the present invention because, for the same number of test compounds, more structure-function relationship information is likely to be revealed.

The method of sensitizing a cell entails selecting a suitable gene or operon. A suitable gene or operon is one whose transcription and/or expression is required for the proliferation of the cell to be sensitized. The next step is to introduce into the cells to be sensitized, an antisense RNA capable of hybridizing to the suitable gene or operon or to the RNA encoded by the suitable gene or operon. Introduction of the antisense RNA can be in the form of a vector in which antisense RNA is produced under the control of an inducible promoter. The amount of antisense RNA produced is modulated by varying an inducer concentration to which the cell is exposed and thereby varying the activity of the promoter driving transcription of the antisense RNA. Thus, cells are sensitized by exposing them to an inducer concentration that results in a sub-lethal level of antisense RNA expression. The requisite maount of inducer may be derived empiracally by one of skill in the art.

In one embodiment of the cell-based assays, antisense nucleic acids complementary to the identified Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi nucleotide sequences or portions thereof (including antisense nucleic acids comprising a nucleotide sequence complementary to one of SEO ID NOs.: 3796-3800, 3806-4860, 5916-10012, and the antisense nucleic acids of SEQ ID NOs.: 8-3795 or antisense nucleic acids comprising a nucleotide sequence complementary to portions of the foregoing nucleic acids thereof), antisense nucleic complementary to homologous coding nucleic acids or portions thereof or homologous antisense nucleic acids are used to inhibit the production of a proliferation-required protein. Vectors producing antisense RNA complementary to identified genes required for proliferation, or portions thereof, are used to limit the concentration of a proliferation-required protein without severely inhibiting growth. The proliferation-required protein may be one of the proteins of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110 or a homologous polypeptide. To achieve that goal, a growth inhibition dose curve of inducer is calculated by plotting various doses of inducer against the corresponding growth inhibition caused by the antisense expression. From this curve, the concentration of inducer needed to achieve various percentages of antisense induced growth inhibition, from 1 to 100% can be determined.

A variety of different regulatable promoters may be used to produce the antisense nucleic acid. Transcription from the regulatable promoters may be modulated by controlling the activity of a transcription factor repressor which acts at the regulatable promoter. For example, if transcription is modulated by affecting the activity of a repressor, the choice of inducer to be used depends on the repressor/operator responsible for regulating transcription of the antisense nucleic acid. If the regulatable promoter comprises a T5 promoter fused to a xylO (xylose operator; e.g. derived from Staphylococcus xylosis (Schnappinger, D. et al., FEMS Microbiol. Let. 129: 121-128 (1995)) then transcription of the antisense nucleic acid may be regulated by a xylose repressor. The xylose

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repressor may be provided by ectoptic expression within an *S. aureus* cell of an exogenous xylose repressor gene, e.g. derived from *S. xylosis* DNA. In such cases transcription of antisense RNA from the promoter is inducible by adding xylose to the medium and the promoter is thus "xylose inducible." Similarly, IPTG inducible promoters may be used. For example, the highest concentration of the inducer that does not reduce the growth rate significantly can be estimated from the curve. Cellular proliferation can be monitored by growth medium turbidity via OD measurements. In another example, the concentration of inducer that reduces growth by 25% can be predicted from the curve. In still another example, a concentration of inducer that reduces growth by 50% can be calculated. Additional parameters such as colony forming units (ofu) can be used to measure cellular viability.

Cells to be assayed are exposed to the above-determined concentrations of inducer. The presence of the inducer at this sub-lethal concentration reduces the amount of the proliferation required gene product to a sub-optimal amount in the cell that will still support growth. Cells grown in the presence of this concentration of inducer are therefore specifically more sensitive to inhibitors of the proliferation-required protein or RNA of interest or to inhibitors of proteins or RNAs in the same biological pathway as the proliferation-required protein or RNA of interest but not to inhibitors of unrelated proteins or RNAs.

Cells pretreated with sub-inhibitory concentrations of inducer and thus containing a reduced amount of proliferation-required target gene product are then used to screen for compounds that reduce cell growth. The sub-lethal concentration of inducer may be any concentration consistent with the intended use of the assay to identify candidate compounds to which the cells are more sensitive. For example, the sub-lethal concentration of the inducer may be such that growth inhibition is at least about 5%, at least about 8%, at least about 10%, at least about 20%, at least about 30%, at least about 40%, at least about 50%, at least about 60% at least about 75%, or more. Cells which are pre-sensitized using the preceding method are more sensitive to inhibitors of the target protein because these cells contain less target protein to inhibit than do wild-type cells.

It will be appreciated that the above cell-based assays may be performed using antisense nucleic acids comprising a nucleotide sequence complementary to any of the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi, or portions thereof, antisense nucleic acids complementary to homologous coding nucleic acids or portions thereof or homologous antisense nucleic acids. In this way, the level or activity of a target, such as any of the proliferation-required polypeptides from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi, or homologous polypeptides.

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In another embodiment of the cell-based assays of the present invention, the level or activity of a proliferation required gene product is reduced using a mutation, such as a temperature sensitive mutation, in the gene encoding a gene product required for proliferation and an antisense nucleic acid comprising a nucleotide sequence complementary to the gene encoding the gene product required for proliferation or a portion thereof. Growing the cells at an intermediate temperature between the permissive and restrictive temperatures of the temperature sensitive mutant where the mutation is in a proliferation-required gene produces cells with reduced activity of the proliferation-required gene product. The antisense RNA complementary to the proliferationrequired sequence further reduces the activity of the proliferation required gene product. Drugs that may not have been found using either the temperature sensitive mutation or the antisense nucleic acid alone may be identified by determining whether cells in which transcription of the antisense nucleic acid has been induced and which are grown at a temperature between the permissive temperature and the restrictive temperature are substantially more sensitive to a test compound than cells in which expression of the antisense nucleic acid has not been induced and which are grown at a permissive temperature. Also drugs found previously from either the antisense nucleic acid alone or the temperature sensitive mutation alone may have a different sensitivity profile when used in cells combining the two approaches, and that sensitivity profile may indicate a more specific action of the drug in inhibiting one or more activities of the gene product.

Temperature sensitive mutations may be located at different sites within the gene and correspond to different domains of the protein. For example, the dnaB gene of Escherichia coli encodes the replication fork DNA helicase. DnaB has several domains, including domains for oligomerization, ATP hydrolysis, DNA binding, interaction with primase, interaction with DnaC, and interaction with DnaA [(Biswas, E.E. and Biswas, S.B. 1999. Mechanism and DnaB helicase of Escherichia coli: structural domains involved in ATP hydrolysis, DNA binding, and oligomerization, Biochem. 38:10919-10928; Hiasa, H. and Marjans, K.J. 1999. Initiation of bidirectional replication at the chromosomal origin is directed by the interaction between helicase and primase, J. Bjol. Chem. 274:27244-27248; San Martin, C., Radermacher, M., Wolpensinger, B., Engel, A., Miles, C.S., Dixon, N.E., and Carazo, J.M. 1998. Three-dimensional reconstructions from cryoelectron microscopy images reveal an intimate complex between helicase DnaB and its loading partner DnaC. Structure 6:501-9; Sutton, M.D., Carr, K.M., Vicente, M., and Kaguni, J.M. 1998. Escherichia coli DnaA protein. The N-terminal domain and loading of DnaB helicase at the E. coli chromosomal origin, J. Biol. Chem. 273:34255-62.)]. Temperature sensitive mutations in different domains of DnaB confer different phenotypes at the restrictive temperature, which include either an abrupt stop or slow stop in DNA replication with or without DNA breakdown (Wechsler, J.A. and Gross, J.D. 1971. Escherichia coli mutants temperature-sensitive for DNA synthesis. Mol. Gen. Genetics 113:273-284) and termination of growth or cell death. Combining the use of temperature sensitive mutations in the dnaB gene that cause cell death at the restrictive temperature

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with an antisense to the *dnaB* gene could lead to the discovery of very specific and effective inhibitors of one or a subset of activities exhibited by DnaB.

It will be appreciated that the above method may be performed with any mutation which reduces but does not eliminate the activity or level of the gene product which is required for proliferation.

It will be appreciated that the above cell-based assays may be performed using mutations in, such as temperature sensitive mutations, and antisense nucleic acids comprising a nucleotide sequence complementary to any of the genes encoding proliferation-required gene products from from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi, or portions thereof (including the nucleic acids of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012), mutations in and antisense nucleic acids complementary to homologous coding nucleic acids or portions thereof or homologous antisense nucleic acids. In this way, the level or activity of a target, such as any of the proliferation-required polypeptides from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi (including the polypeptides of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110), or homologous polypeptides may be reduced.

When screening for antimicrobial agents against a gene product required for proliferation, growth inhibition of cells containing a limiting amount of that proliferation-required gene product can be assayed. Growth inhibition can be measured by directly comparing the amount of growth, measured by the optical density of the growth medium, between an experimental sample and a control sample. Alternative methods for assaying cell proliferation include measuring green fluorescent protein (GFP) reporter construct emissions, various enzymatic activity assays, and other methods well known in the art.

It will be appreciated that the above method may be performed in solid phase, liquid phase or a combination of the two. For example, cells grown on nutrient agar containing the inducer of the antisense construct may be exposed to compounds spotted onto the agar surface. If desired, the cells may be grown on agar containing varying concentrations of the inducer. A compound's effect may be judged from the diameter of the resulting killing zone, the area around the compound application point in which cells do not grow. Multiple compounds may be transferred to agar plates and simultaneously tested using automated and semi-automated equipment including but not restricted to multi-channel pipettes (for example the Beckman Multimek) and multi-channel spotters (for example the Genomic Solutions Flexys). In this way multiple plates and thousands to millions of compounds may be tested per day.

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The compounds may also be tested entirely in liquid phase using microtiter plates as described below. Liquid phase screening may be performed in microtiter plates containing 96, 384, 1536 or more wells per microtiter plate to screen multiple plates and thousands to millions of compounds per day. Automated and semi-automated equipment may be used for addition of reagents (for example cells and compounds) and determination of cell density.

EXAMPLE 9

Cell-Based Assay Using Antisense Complementary to Genes Encoding Ribosomal Proteins

The effectiveness of the above cell-based assay was validated using constructs transribing antisense RNA to the proliferation required E. coli genes rplL, rplJ, and rplW encoding ribosomal proteins L7/L12, L10 and L23 respectively. These proteins are essential components of the protein synthesis apparatus of the cell and as such are required for proliferation. These constructs were used to test the effect of antisense transcription on cell sensitivity to antibiotics known to bind to the ribosome and thereby inhibit protein synthesis. Constructs transcribing antisense RNA to several other genes (elaD, visC, yohH, and atpE/B), the products of which are not involved in protein synthesis were used for comparison.

First, pLex5BA (Krause et al., J. Mol. Biol. 274: 365 (1997)) vectors containing antisense constructs to either *rplW* or to *elaD* were introduced into separate *E. coli* cell populations. Vector introduction is a technique well known to those of ordinary skill in the art. The vectors of this example contain IPTG inducible promoters that drive the transcription of the antisense RNA in the presence of the inducer. However, those skilled in the art will appreciate that other inducible promoters may also be used. Suitable vectors are also well known in the art. Antisense clones to genes encoding different ribosomal proteins or to genes encoding proteins that are not involved in protein synthesis were utilized to test the effect of antisense transcription on cell sensitivity to the antibiotics known to bind to ribosomal proteins and inhibit protein synthesis. Antisense nucleic acids comprising a nucleotide sequence complementarty to the *elaD*, *atpB&atpE*, *visC* and *yohH* genes are referred to as AS-*elaD*, AS-*atpB/E*, AS-*visC*, AS-*yohH* respectively. These genes are not known to be involved in protein synthesis. Antisense nucleic acids to the *rplL*, *rplL&rplJ* and *rplW* genes are referred to as AS-*rplL*, AS-*rplL/J*, and AS-*rplW* respectively. These genes encode ribosomal proteins L7/L12 (*rplL*) L10 (*rplJ*) and L23 (*rplW*). Vectors containing these antisense nucleic acids were introduced into separate *E. coli* cell populations.

The cell populations containing vectors producing AS-elaD or AS-rplW were exposed to a range of IPTG concentrations in liquid medium to obtain the growth inhibitory dose curve for each clone (Fig. 1). First, seed cultures were grown to a particular turbidity measured by the optical density (OD) of the growth solution. The OD of the solution is directly related to the number of bacterial cells contained therein. Subsequently, sixteen 200 µl liquid medium cultures were grown in a 96 well microtiter plate at 37° C with a range of IPTG concentrations in duplicate two-fold

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serial dilutions from 1600 uM to 12.5 µM (final concentration). Additionally, control cells were grown in duplicate without IPTG. These cultures were started from an inoculum of equal amounts of cells derived from the same initial seed culture of a clone of interest. The cells were grown for up to 15 hours and the extent of growth was determined by measuring the optical density of the cultures at 600 nm. When the control culture reached mid-log phase the percent growth (relative to the control culture) for each of the IPTG containing cultures was plotted against the log concentrations of IPTG to produce a growth inhibitory dose response curve for the IPTG. The concentration of IPTG that inhibits cell growth to 50% (IC₅₀) as compared to the 0 mM IPTG control (0% growth inhibition) was then calculated from the curve. Under these conditions, an amount of antisense RNA was produced that reduced the expression levels of *rplW* or *elaD* to a degree such that growth of cells containing their respective antisense vectors was inhibited by 50%.

Alternative methods of measuring growth are also contemplated. Examples of these methods include measurements of proteins, the expression of which is engineered into the cells being tested and can readily be measured. Examples of such proteins include green fluorescent protein (GFP), luciferase, and various enzymes.

Cells were pretreated with the selected concentration of IPTG and then used to test the sensitivity of cell populations to tetracycline, erythromycin and other known protein synthesis inhibitors. Figure 1 is an IPTG dose response curve in *E. coli* transformed with an IPTG-inducible plasmid containing either an antisense clone to the *E. coli rplW* gene (AS-*rplW*) which encodes ribosomal protein L23 which is required for protein synthesis and essential for cell proliferation, or an antisense clone to the *elaD* (AS-*elaD*) gene which is not known to be involved in protein synthesis.

An example of a tetracycline dose response curve is shown in Figures 2A and 2B for the rplW and elaD genes, respectively. Cells were grown to log phase and then diluted into medium alone or medium containing IPTG at concentrations which give 20% and 50% growth inhibition as determined by IPTG dose response curves. After 2.5 hours, the cells were diluted to a final OD₆₀₀ of 0.002 into 96 well plates containing (1) +/- IPTG at the same concentrations used for the 2.5 hour pre-incubation; and (2) serial two-fold dilutions of tetracycline such that the final concentrations of tetracycline range from 1 μ g/ml to 15.6 ng/ml and 0 μ g/ml. The 96 well plates were incubated at 37°C and the OD₆₀₀ was read by a plate reader every 5 minutes for up to 15 hours. For each IPTG concentration and the no IPTG control, tetracycline dose response curves were determined when the control (absence of tetracycline) reached 0.1 OD₆₀₀.

To compare tetracycline sensitivity with and without IPTG, tetracycline IC_{50s} were determined from the dose response curves (Figs. 3A-B). Cells transcribing antisense nucleic acids AS-rplL or AS-rplW to genes encoding ribosomal proteins L7/L12 and L23 respectively showed increased sensitivity to tetracycline (Fig. 2A) as compared to cells with reduced levels of the elaD

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gene product (AS-elaD) (Fig. 2B). Figure 3 shows a summary bar chart in which the ratios of tetracycline IC_{50s} determined in the presence of IPTG which gives 50% growth inhibition versus tetracycline IC_{50s} determined without IPTG (fold increase in tetracycline sensitivity) were plotted. Cells with reduced levels of either L7/L12 (encoded by genes rplL, rplJ) or L23 (encoded by the rplW gene) showed increased sensitivity to tetracycline (Fig. 3). Cells expressing antisense to genes not known to be involved in protein synthesis (AS-atpB/E, AS-visC, AS-elaD, AS-yohH) did not show the same increased sensitivity to tetracycline, validating the specificity of this assay (Fig. 3).

In addition to the above, it has been observed in initial experiments that clones transcribing antisense RNA to genes involved in protein synthesis (including genes encoding ribosomal proteins L7/L12 & L10, L7/L12 alone, L22, and L18, as well as genes encoding rRNA and Elongation Factor G) have increased sensitivity to the macrolide, erythromycin, whereas clones transcribing antisense to the non-protein synthesis genes *elaD*, *atpB/E* and *visC* do not. Furthermore, the clone transcribing antisense to *rplL* and *rplJ* (AS-*rplL/J*) does not show increased sensitivity to nalidixic acid and ofloxacin, antibiotics which do not inhibit protein synthesis.

The results with the ribosomal protein genes rplL, rplJ, and rplW as well as the initial results using various other antisense clones and antibiotics show that limiting the concentration of an antibiotic target makes cells more sensitive to the antimicrobial agents that specifically interact with that protein. The results also show that these cells are sensitized to antimicrobial agents that inhibit the overall function in which the protein target is involved but are not sensitized to antimicrobial agents that inhibit other functions. It will be appreciated that the cell-based assays described above may be implemented using the Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi antisense nucleotide sequences which inhibit the activity of genes required for proliferation described herein (including the antisense nucleic acids of SEQ ID NOs.: 8-3795) or antisense nucleic acids comprising nucleotide sequences which are complementary to the sequences of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012 or portions thereof.

It will be appreciated that the above cell-based assays may be performed using antisense nucleic acids complementary to any of the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi, or portions thereof, antisense nucleic acids complementary to homologous coding nucleic acids or portions thereof, or homologous antisense nucleic acids. In this way, the level or activity of a target, such as any of the proliferation-required polypeptides from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa,

Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi, or homologous polypeptides may be reduced.

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The cell-based assay described above may also be used to identify the biological pathway in which a proliferation-required nucleic acid or its gene product lies. In such methods, cells transcribing a sub-lethal level of antisense to a target proliferation-required nucleic acid and control cells in which transcription of the antisense has not been induced are contacted with a panel of antibiotics known to act in various pathways. If the antibiotic acts in the pathway in which the target proliferation-required nucleic acid or its gene product lies, cells in which transcription of the antisense has been induced will be more sensitive to the antibiotic than cells in which expression of the antisense has not been induced.

As a control, the results of the assay may be confirmed by contacting a panel of cells transcribing antisense nucleic acids to many different proliferation-required genes including the target proliferation-required gene. If the antibiotic is acting specifically, heightened sensitivity to the antibiotic will be observed only in the cells transcribing antisense to a target proliferation-required gene (or cells expressing antisense to other proliferation-required genes in the same pathway as the target proliferation-required gene) but will not be observed generally in all cells expressing antisense to proliferation-required genes.

It will be appreciated that the above cell-based assays may be performed using antisense nucleic acids complementary to any of the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi, (including antisense nucleic acids complementary to SEQ ID NOs: 3796-3800, 3806-4860, 5916-10012, or the antisense nucleic acids of SEQ ID NOs.: 8-3795) or portions thereof, antisense nucleic acids comprising nucleotide sequences complementary to homologous coding nucleic acids or portions thereof, or homologous antisense nucleic acids In this way, the level or activity of a target, such as any of the proliferation-required polypeptides from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi (including the polypeptides of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110), or homologous polypeptides may be reduced.

Similarly, the above method may be used to determine the pathway on which a test compound, such as a test antibiotic acts. A panel of cells, each of which transcribes an antisense to a proliferation-required nucleic acid in a known pathway, is contacted with a compound for which it is desired to determine the pathway on which it acts. The sensitivity of the panel of cells to the test compound is determined in cells in which transcription of the antisense has been induced and in control cells in which expression of the antisense has not been induced. If the test compound acts on the pathway on which an antisense nucleic acid acts, cells in which expression of the antisense

has been induced will be more sensitive to the compound than cells in which expression of the antisense has not been induced. In addition, control cells in which expression of antisense to proliferation-required genes in other pathways has been induced will not exhibit heightened sensitivity to the compound. In this way, the pathway on which the test compound acts may be determined.

It will be appreciated that the above cell-based assays may be performed using antisense nucleic acids comprising nucleotide sequences complementary to any of the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi (including antisense nucleic acids complementary to SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, such as the antisense nucleic acids of SEQ ID NOs.: 8-3795) or portions thereof, antisense nucleic acids complementary to homologous coding nucleic acids or portions thereof, or homologous antisense nucleic acids In this way, the level or activity of a target, such as any of the proliferation-required polypeptides from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi (including the polypeptides of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110) or homologous polypeptides may be reduced.

The Example below provides one method for performing such assays.

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EXAMPLE 10

Identification of the Pathway in which a Proliferation-Required

Gene Lies or the Pathway on which an Antibiotic Acts

A. Preparation of Bacterial Stocks for Assav

To provide a consistent source of cells to screen, frozen stocks of host bacteria containing the desired antisense construct are prepared using standard microbiological techniques. For example, a single clone of the microorganism can be isolated by streaking out a sample of the original stock onto an agar plate containing nutrients for cell growth and an antibiotic for which the antisense construct contains a selectable marker which confers resistance. After overnight growth an isolated colony is picked from the plate with a sterile needle and transferred to an appropriate liquid growth medium containing the antibiotic required for maintenance of the plasmid. The cells are incubated at 30°C to 37°C with vigorous shaking for 4 to 6 hours to yield a culture in exponential growth. Sterile glycerol is added to 15% (volume to volume) and 100μL to 500 μL aliquots are distributed into sterile cryotubes, snap frozen in liquid nitrogen, and stored at -80°C for future assays.

B. Growth of Bacteria for Use in the Assay

A day prior to an assay, a stock vial is removed from the freezer, rapidly thawed (37°C water bath) and a loop of culture is streaked out on an agar plate containing nutrients for cell growth and an antibiotic to which the selectable marker of the antisense construct confers resistance. After overnight growth at 37°C, ten randomly chosen, isolated colonies are transferred from the plate (sterile inoculum loop) to a sterile tube containing 5 mL of LB medium containing the antibiotic to which the antisense vector confers resistance. After vigorous mixing to form a homogeneous cell suspension, the optical density of the suspension is measured at 600 nm (OD₆₀₀) and if necessary an aliquot of the suspension is diluted into a second tube of 5 mL, sterile, LB medium plus antibiotic to achieve an OD₆₀₀ \leq 0.02 absorbance units. The culture is then incubated at 37° C for 1-2 hrs with shaking until the OD₆₀₀ reaches OD 0.2 – 0.3. At this point the cells are ready to be used in the assay.

C. Selection of Media to be Used in Assay

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Two-fold dilution series of the inducer are generated in culture media containing the appropriate antibiotic for maintenance of the antisense construct. Several media are tested side by side and three to four wells are used to evaluate the effects of the inducer at each concentration in each media. For example, LB broth, TBD broth and Muller-Hinton media may be tested with the inducer xylose at the following concentrations, 5 mM, 10 mM, 20 mM, 40 mM, 80 mM, 120 mM and 160 mM. Equal volumes of test media-inducer and cells are added to the wells of a 384 well microtiter plate and mixed. The cells are prepared as described above and diluted 1:100 in the appropriate media containing the test antibiotic immediately prior to addition to the microtiter plate wells. For a control, cells are also added to several wells of each media that do not contain inducer, for example 0 mM xylose. Cell growth is monitored continuously by incubation at 37°C in a microtiter plate reader monitoring the OD600 of the wells over an 18-hour period. The percent inhibition of growth produced by each concentration of inducer is calculated by comparing the rates of logarithmic growth against that exhibited by cells growing in medium without inducer. The medium yielding greatest sensitivity to inducer is selected for use in the assays described below.

D. Measurement of Test Antibiotic Sensitivity in the Absence of Antisense Construct Induction

Two-fold dilution series of antibiotics of known mechanism of action are generated in the culture medium selected for further assay development that has been supplemented with the antibiotic used to maintain the construct. A panel of test antibiotics known to act on different pathways is tested side by side with three to four wells being used to evaluate the effect of a test antibiotic on cell growth at each concentration. Equal volumes of test antibiotic and cells are added to the wells of a 384 well microtiter plate and mixed. Cells are prepared as described above using the medium selected for assay development supplemented with the antibiotic required to maintain the antisense construct and are diluted 1:100 in identical medium immediately prior to addition to the microtiter plate wells. For a control, cells are also added to several wells that lack antibiotic,

but contain the solvent used to dissolve the antibiotics. Cell growth is monitored continuously by incubation at 37° C in a microtiter plate reader monitoring the OD_{600} of the wells over an 18-hour period. The percent inhibition of growth produced by each concentration of antibiotic is calculated by comparing the rates of logarithmic growth against that exhibited by cells growing in medium without antibiotic. A plot of percent inhibition against log[antibiotic concentration] allows extrapolation of an IC_{50} value for each antibiotic.

E. Measurement of Test Antibiotic Sensitivity in the Presence of Antisense Construct Inducer

The culture medium selected for use in the assay is supplemented with inducer at concentrations shown to inhibit cell growth by 50% and 80% as described above, as well as the antibiotic used to maintain the construct. Two-fold dilution series of the panel of test antibiotics used above are generated in each of these media. Several antibiotics are tested side by side in each medium with three to four wells being used to evaluate the effects of an antibiotic on cell growth at each concentration. Equal volumes of test antibiotic and cells are added to the wells of a 384 well microtiter plate and mixed. Cells are prepared as described above using the medium selected for use in the assay supplemented with the antibiotic required to maintain the antisense construct. The cells are diluted 1:100 into two 50 mL aliquots of identical medium containing concentrations of inducer that have been shown to inhibit cell growth by 50% and 80 % respectively and incubated at 37°C with shaking for 2.5 hours. Immediately prior to addition to the microtiter plate wells, the cultures are adjusted to an appropriate OD₆₀₀ (typically 0.002) by dilution into warm (37°C) sterile medium supplemented with identical concentrations of the inducer and antibiotic used to maintain the antisense construct. For a control, cells are also added to several wells that contain solvent used to dissolve test antibiotics but which contain no antibiotic. Cell growth is monitored continuously by incubation at 37°C in a microtiter plate reader monitoring the OD₆₀₀ of the wells over an 18-hour period. The percent inhibition of growth produced by each concentration of antibiotic is calculated by comparing the rates of logarithmic growth against that exhibited by cells growing in medium without antibiotic. A plot of percent inhibition against log[antibiotic concentration] allows extrapolation of an IC₅₀ value for each antibiotic.

F. Determining the Specificity of the Test Antibiotics

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A comparison of the IC₅₀s generated by antibiotics of known mechanism of action under antisense induced and non-induced conditions allows the pathway in which a proliferation-required nucleic acid lies to be identified. If cells expressing an antisense nucleic acid comprising a nucleotide sequence complementary to a proliferation-required gene are selectively sensitive to an antibiotic acting via a particular pathway, then the gene against which the antisense acts is involved in the pathway on which the antibiotic acts.

G. Identification of Pathway in which a Test Antibiotic Acts

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As discussed above, the cell-based assay may also be used to determine the pathway against which a test antibiotic acts. In such an analysis, the pathways against which each member of a panel of antisense nucleic acids acts are identified as described above. A panel of cells, each containing an inducible vector which transcribes an antisense nucleic acid comprising a nucleotide sequence complementary to a gene in a known proliferation-required pathway, is contacted with a test antibiotic for which it is desired to determine the pathway on which it acts under inducing and non-inducing conditions. If heightened sensitivity is observed in induced cells transcribing antisense complementary to a gene in a particular pathway but not in induced cells transcribing antisense nucleic acids comprising nucleotide sequences complementary to genes in other pathways, then the test antibiotic acts against the pathway for which heightened sensitivity was observed.

One skilled in the art will appreciate that further optimization of the assay conditions, such as the concentration of inducer used to induce antisense transcription and/or the growth conditions used for the assay (for example incubation temperature and medium components) may further increase the selectivity and/or magnitude of the antibiotic sensitization exhibited.

It will be appreciated that the above cell-based assays may be performed using antisense nucleic acids comprising nucleotide sequences complementary to any of the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi, (including antisense nucleic acids comprising nucleotide sequences complemenatary to SEQ ID NOs.: 8-3795) or portions thereof, 5916-10012, such as the antisense nucleic acids of SEQ ID NOs.: 8-3795) or portions thereof, antisense nucleic acids complementary to homologous coding nucleic acids or portions thereof, or homologous antisense nucleic acids In this way, the level or activity of a target, such as any of the proliferation-required polypeptides from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi (including the polypeptides of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110), or homologous polypeptides may be reduced.

The following example confirms the effectiveness of the methods described above.

EXAMPLE 11 <u>Identification of the Biological Pathway in which a Proliferation-Required Gene Lies</u>

The effectiveness of the above assays was validated using proliferation-required genes from *E. coli* which were identified using procedures similar to those described above. Antibiotics of various chemical classes and modes of action were purchased from Sigma Chemicals (St. Louis, MO). Stock solutions were prepared by dissolving each antibiotic in an appropriate aqueous

solution based on information provided by the manufacturer. The final working solution of each

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antibiotic contained no more than 0.2% (w/v) of any organic solvent. To determine their potency against a bacterial strain engineered for transcription of an antisense comprising a nucleotide sequence complementary to a proliferation-required 50S ribosomal protein, each antibiotic was serially diluted two- or three- fold in growth medium supplemented with the appropriate antibiotic for maintenance of the antisense construct. At least ten dilutions were prepared for each antibiotic. 25 µL aliquots of each dilution were transferred to discrete wells of a 384-well microplate (the assay plate) using a multi-channel pipette. Quadruplicate wells were used for each dilution of an antibiotic under each treatment condition (plus and minus inducer). Each assay plate contained twenty wells for cell growth controls (growth medium replacing antibiotic), ten wells for each treatment (plus and minus inducer, in this example IPTG). Assay plates were usually divided into the two treatments: half the plate containing induced cells and an appropriate concentrations of inducer (in this example IPTG) to maintain the state of induction, the other half containing non-induced cells in the absence of IPTG.

Cells for the assay were prepared as follows. Bacterial cells containing a construct, from which transcription of antisense nucleic acid comprising a nucleotide sequence complementary to rplL and rplJ (AS-rplL/J), which encode proliferation-required 50S ribosomal subunit proteins, is inducible in the presence of IPTG, were grown into exponential growth (OD600 0.2 to 0.3) and then diluted 1:100 into fresh medium containing either 400 µM or 0 µM inducer (IPTG). These cultures were incubated at 37° C for 2.5 hr. After a 2.5 hr incubation, induced and non-induced cells were respectively diluted into an assay medium at a final OD600 value of 0.0004. The medium contained an appropriate concentration of the antibiotic for the maintenance of the antisense construct. In addition, the medium used to dilute induced cells was supplemented with 800 µM IPTG so that addition to the assay plate would result in a final IPTG concentration of 400 µM. Induced and noninduced cell suspensions were dispensed (25 µl/well) into the appropriate wells of the assay plate as discussed previously. The plate was then loaded into a plate reader, incubated at constant temperature, and cell growth was monitored in each well by the measurement of light scattering at 595 nm. Growth was monitored every 5 minutes until the cell culture attained a stationary growth phase. For each concentration of antibiotic, a percentage inhibition of growth was calculated at the time point corresponding to mid-exponential growth for the associated control wells (no antibiotic, plus or minus IPTG). For each antibiotic and condition (plus or minus IPTG), a plot of percent inhibition versus log of antibiotic concentration was generated and the IC50 determined. A comparison of the IC₅₀ for each antibiotic in the presence and absence of IPTG revealed whether induction of the antisense construct sensitized the cell to the mechanism of action exhibited by the antibiotic. Cells which exhibited a statistically significant decrease in the IC₅₀ value in the presence of inducer were considered to have an increased sensitivity to the test antibiotic.

The results are provided in the table below, which lists the classes and names of the antibiotics used in the analysis, the targets of the antibiotics, the IC_{50} in the absence of IPTG, the IC_{50} in the presence of IPTG, the concentration units for the IC_{50} s, the fold increase in IC_{50} in the presence of IPTG, and whether increased sensitivity was observed in the presence of IPTG.

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<u>TABLE III</u> et of Expression of Antisense RNA to rplL and rplJ on Antibiotic Sensi

Effect of Ex	Effect of Expression of Antisense RNA to rplL and rplJ on Antibiotic Sensitivity	and rplJ on	Antibiotic Sens	itivity		
ANTIBIOTIC CLASS /Names	TARGET	IC _{so} (-IPTG)	IC ₅₀ (+IPTG)	Cone. Unit	Fold Increase in Sensitivity	Sensitivity Increased?
PROTEIN SYNTHESIS INHIBITOR						
AMINOGLYCOSIDES				-		
Gentamicin	30S ribosome function	2715	19.19	ng/ml	141	Yes
Streptomycin	30S ribosome function	11280	191	lm/gu	92	Yes
Spectinomycin	30S ribosome function	18050	<156	lm/gu		Yes
Tobramycin	30S ribosome function	3594	70.58	lm/gu	51	Yes
MACROLIDES						
Erythromycin	50S ribosome function	7467	187	lm/gu	40	Yes
AROMATIC POYKETIDES			-			
Tetracycline	30S ribosome function	199.7	1.83	ng/ml	109	Yes
Minocycline	30S ribosome function	668.4	3.897	lm/gu	172	Yes
Doxycycline	30S ribosome function	413.1	27.81	ng/ml	15	Yes
OTHER PROTEIN SYNTHESIS INHIBITORS		٠				
Fusidic acid	Elongation Factor G function	29990	641	lm/gu	94	Yes
Chloramphenicol	30S ribosome function	465.4	1.516	ng/ml	307	Yes
Lincomycin	50S ribosome function	47150	324.2	ng/ml	145	Yes
OTHER ANTIBIOTIC MECHANISMS						
B-LACTAMS						
Cefoxitin	Cell wall biosynthesis	2782	2484	lm/gu	_	N _o
Cefotaxime	Cell wall biosynthesis	24.3	24.16	ng/ml	1	No
DNA SYNTHESIS INHIBITORS						
Nalidixic acid	DNA Gyrase activity	6973	6025	ng/ml	_	No
Offoxacin	DNA Gyrase activity	49.61	45.89	lm/gu	1	No
OTHER						
Bacitracin	Cell membrane function	4077	4677	mg/ml	1	No
Trimethoprim	Dihydrofolate Reductase activity	128.9	181.97	ng/ml	_	% %
Vancomycin	Cell wall biosynthesis	145400	72550	ng/ml	2	No No

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The above results demonstrate that induction of an antisense RNA complementary to genes encoding 50S ribosomal subunit proteins results in a selective and highly significant sensitization of cells to antibiotics that inhibit ribosomal function and protein synthesis. The above results further demonstrate that induction of an antisense to an essential gene sensitizes a cell or microorganism to compounds that interfere with that gene product's biological role. This sensitization is restricted to compounds that interfere with pathways associated with the targeted gene and its product.

It will be appreciated that the above cell-based assays may be performed using antisense nucleic acids complementary to any of the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi (including antisense nucleic acids complementary to SEQ ID NOs.: 8-3795) or portions thereof, antisense nucleic acids complementary to homologous coding nucleic acids or portions thereof or homologous antisense nucleic acids. In this way, the level or activity of a target, such as any of the proliferation-required polypeptides from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi i (including the polypeptides of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110), or homologous polypeptides may be reduced.

Example 11A below describes an analysis performed in Staphylococcus aureus.

EXAMPLE 11A

Identification of the Biological Pathway in which a Gene Required for Proliferation of Staphylococcus aureus Lies

Antibiotics of various chemical classes and modes of action were purchased from chemical suppliers, for example Sigma Chemicals (St. Louis, MO). Stock solutions were prepared by dissolving each antibiotic in an appropriate aqueous solution based on information provided by the manufacturer. The final working solution of each antibiotic contained no more than 0.2% (w/v) of any organic solvent.

To determine its potency against a bacterial strain containing an antisense nucleic acid comprising a nucleotide sequence complementary to the nucleotide sequence encoding the Beta subunit of DNA gyrase (which is required for proliferation) under the control of a xylose inducible promoter, each antibiotic was serially diluted two- or three- fold in growth medium supplemented with the appropriate antibiotic for maintenance of the antisense construct. At least ten dilutions were prepared for each antibiotic.

Aliquots (25 μ L) of each dilution were transferred to discrete wells of a 384-well microplate (the assay plate) using a multi-channel pipette. Quadruplicate wells were used for each dilution of an antibiotic under each treatment condition (plus and minus inducer). Each assay plate

contained twenty wells for cell growth controls (growth medium, no antibiotic), ten wells for each treatment (plus and minus inducer, xylose, in this example). Half the assay plate contained induced cells (in this example *Staphylococcus aureus* cells) and appropriate concentrations of inducer (xylose, in this example) to maintain the state of induction while the other half of the assay plate contained non-induced cells maintained in the absence of inducer.

Preparation of Bacterial Cells

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Cells of a bacterial clone containing a construct in which transcription of antisense comprising a nucleotide sequence complementary to the sequence encoding the Beta subunit of DNA gyrase under the control of the xylose inducible promoter (S1M10000001F08) were grown into exponential growth (OD₆₀₀ 0.2 to 0.3) and then diluted 1:100 into fresh medium containing either 12 mM or 0 mM inducer (xylose). These cultures were incubated at 37° C for 2.5 hr. The presence of inducer (xylose) in the medium initiates and maintains production of antisense RNA from the antisense construct. After a 2.5 hr incubation, induced and non-induced cells were respectively diluted into an assay medium containing an appropriate concentration of the antibiotic for the maintenance of the antisense construct. In addition, medium used to dilute induced cells was supplemented with 24 mM xylose so that addition to the assay plate would result in a final xylose concentration of 12 mM. The cells were diluted to a final OD₆₀₀ value of 0.0004.

Induced and non-induced cell suspensions were dispensed (25 µl/well) into the appropriate wells of the assay plate as discussed previously. The plate was then loaded into a plate reader and incubated at constant temperature while cell growth was monitored in each well by the measurement of light scattering at 595 nm. Growth was monitored every 5 minutes until the cell culture attained a stationary growth phase. For each concentration of antibiotic, a percentage inhibition of growth was calculated at the time point corresponding to mid-exponential growth for the associated control wells (no antibiotic, plus or minus xylose). For each antibiotic and condition (plus or minus xylose), plots of percent inhibition versus Log of antibiotic concentration were generated and IC_{50s} determined.

A comparison of each antibiotic's IC_{50} in the presence and absence of inducer (xylose, in this example) reveals whether induction of the antisense construct sensitized the cell to the antibiotic's mechanism of action. If the antibiotic acts against the β subunit of DNA gyrase, the IC_{50} of induced cells will be significantly lower than the IC_{50} of uninduced cells.

Figure 4 lists the antibiotics tested, their targets, and their fold increase in potency between induced cells and uninduced cells. As illustrated in Figure 4, the potency of cefotaxime, cefoxitin, fusidic acid, lincomycin, tobramycin, trimethoprim and vancomycin, each of which act on targets other than the β subunit of gyrase, was not significantly different in induced cells as compared to uninduced cells. However, the potency of novobiocin, which is known to act against the Beta subunit of DNA gyrase, was significantly different between induced cells and uninduced cells.

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Thus, induction of an antisense nucleic acid comprising a nucleotide sequence complementary to the sequence encoding the β subunit of gyrase results in a selective and significant sensitization of *Staphylococcus aureus* cells to an antibiotic which inhibits the activity of this protein. Furthermore, the results demonstrate that induction of an antisense construct to an essential gene sensitizes a cell or microorganism to compounds that interfere with that gene product's biological role. This sensitization is apparently restricted to compounds that interfere with the targeted gene and its product.

It will be appreciated that the above cell-based assays may be performed using antisense nucleic acids complementary to any of the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi (including antisense nucleic acids complementary to SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, such as the antisense nucleic acids of SEQ ID NOs. 8-3795), or portions thereof, antisense nucleic acids complementary to homologous coding nucleic acids or portions thereof, or homologous antisense nucleic acids. In this way, the level or activity of a target, such as any of the proliferation-required polypeptides from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi, or homologous polypeptides may be reduced.

Assays utilizing antisense constructs to essential genes or portions thereof can be used to identify compounds that interfere with the activity of those gene products. Such assays could be used to identify drug leads, for example antibiotics.

Panels of cells transcribing different antisense nucleic acids can be used to characterize the point of intervention of a compound affecting an essential biochemical pathway including antibiotics with no known mechanism of action.

Assays utilizing antisense constructs to essential genes can be used to identify compounds that specifically interfere with the activity of multiple targets in a pathway. Such constructs can be used to simultaneously screen a sample against multiple targets in one pathway in one reaction (Combinatorial HTS).

Furthermore, as discussed above, panels of antisense construct-containing cells may be used to characterize the point of intervention of any compound affecting an essential biological pathway including antibiotics with no known mechanism of action.

It will be appreciated that the above cell-based assays may be performed using antisense nucleic acids complementary to any of the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi (including antisense nucleic acids comprising nucleotide sequences

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complementary to SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, such as the antisense nucleic acids of SEQ ID NOs. 8-3795), or portions thereof, antisense nucleic acids complementary to homologous coding nucleic acids or portions thereof, or homologous antisense nucleic acids. In this way, the level or activity of a target, such as any of the proliferation-required polypeptides from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi or homologous polypeptides may be reduced.

Another embodiment of the present invention is a method for determining the pathway against which a test antibiotic compound is active, in which the activity of target proteins or nucleic acids involved in proliferation-required pathways is reduced by contacting cells with a sub-lethal concentration of a known antibiotic which acts against the target protein or nucleic acid. In one embodiment, the target protein or nucleic acid corresponds to a proliferation-required nucleic acid identified using the methods described above, such as the polypeptides of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110, or homologous polypeptides. The method is similar to those described above for determining which pathway a test antibiotic acts against, except that rather than reducing the activity or level of a proliferation-required gene product using a sub-lethal level of antisense to a proliferation-required gene product using a sub-lethal level of a known antibiotic which acts against the proliferation required gene product. Heightened sensitivity determines the pathway on which the test compound is active.

Interactions between drugs which affect the same biological pathway have been described in the literature. For example, Mecillinam (Amdinocillin) binds to and inactivates the penicillin binding protein 2 (PBP2, product of the mrdA in E. coli). This antibiotic interacts with other antibiotics that inhibit PBP2 as well as antibiotics that inhibit other penicillin binding proteins such as PBP3 [(Gutmann, L., Vincent, S., Billot-Klein, D., Acar, J.F., Mrena, E., and Williamson, R. (1986) Involvement of penicillin-binding protein 2 with other penicillin-binding proteins in lysis of Escherichia coli by some beta-lactam antibiotics alone and in synergistic lytic effect of amdinocillin (mecillinam). Antimicrobial Agents & Chemotherapy, 30:906-912]]. Interactions between drugs could, therefore, involve two drugs that inhibit the same target protein or nucleic acid or inhibit different proteins or nucleic acids in the same pathway [(Fukuoka, T., Domon, H., Kakuta, M., Ishii, C., Hirasawa, A., Utsui, Y., Ohya, S., and Yasuda, H. (1997) Combination effect between panipenem and vancomycin on highly methicillin-resistant Staphylococcus aureus. Japan. J. Antibio. 50:411-419; Smith, C.E., Foleno, B.E., Barrett, J.F., and Frosc, M.B. (1997) Assessment of the synergistic interactions of levofloxacin and ampicillin against Enterococcus faecium by the checkerboard agar dilution and time-kill methods. Diagnos. Microbiol. Infect. Disease 27:85-92; den Hollander, J.G., Horrevorts, A.M., van Goor, M.L., Verbrugh, H.A., and Mouton, J.W. (1997)

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Synergism between tobramycin and ceftazidime against a resistant *Pseudomonas aeruginosa* strain, tested in an in vitro pharmacokinetic model. Antimicrobial Agents & Chemotherapy, 41:95-110)].

Two drugs may interact even though they inhibit different targets. For example, the proton pump inhibitor, Omeprazole, and the antibiotic, Amoxycillin, two synergistic compounds acting together, can cure *Helicobacter pylori* infection [(Gabryelewicz, A., Laszewicz, W., Dzieniszewski, J., Ciok, J., Marlicz, K., Bielecki, D., Popiela, T., Legutko, J., Knapik, Z., Poniewierka, E. (1997) Multicenter evaluation of dual-therapy (omeprazol and amoxycillin) for *Helicobacter pylori*-associated duodenal and gastric ulcer (two years of the observation). J. Physiol. Pharmacol. 48 Suppl 4:93-105)].

The growth inhibition from the sub-lethal concentration of the known antibiotic may be at least about 5%, at least about 8%, at least about 10%, at least about 20%, at least about 30%, at least about 40%, at least about 50%, at least about 60%, or at least about 75%, or more.

Alternatively, the sub-lethal concentration of the known antibiotic may be determined by measuring the activity of the target proliferation-required gene product rather than by measuring growth inhibition.

Cells are contacted with a combination of each member of a panel of known antibiotics at a sub-lethal level and varying concentrations of the test antibiotic. As a control, the cells are contacted with varying concentrations of the test antibiotic alone. The IC_{50} of the test antibiotic in the presence and absence of the known antibiotic is determined. If the IC_{50} s in the presence and absence of the known drug are substantially similar, then the test drug and the known drug act on different pathways. If the IC_{50} s are substantially different, then the test drug and the known drug act on the same pathway.

It will be appreciated that the above cell-based assays may be performed using a sub-lethal concentration of a known antibiotic which acts against the product of any of the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi (including the products of SEQ ID NOs: 3796-3800, 3806-4860, 5916-10012, or portions thereof, or the products of homologous coding nucleic acids or portions thereof. In this way, the level or activity of a target, such as any of the proliferation-required polypeptides from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi (including the polypeptides of SEQ ID NOs.: 3801-3805, 4861-5915, 10013-14110), or homologous polypeptides may be reduced.

Another embodiment of the present invention is a method for identifying a candidate compound for use as an antibiotic in which the activity of target proteins or nucleic acids involved in proliferation-required pathways is reduced by contacting cells with a sub-lethal concentration of

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a known antibiotic which acts against the target protein or nucleic acid. In one embodiment, the target protein or nucleic acid is a target protein or nucleic acid corresponding to a proliferation-required nucleic acid identified using the methods described above. The method is similar to those described previously herein for identifying candidate compounds for use as antibiotics except that rather than reducing the activity or level of a proliferation-required gene product using a sub-lethal level of antisense to a proliferation-required nucleic acid, the activity or level of the proliferation-required gene product is reduced using a sub-lethal level of a known antibiotic which acts against the proliferation required gene product.

The growth inhibition from the sub-lethal concentration of the known antibiotic may be at least about 5%, at least about 8%, at least about 10%, at least about 20%, at least about 30%, at least about 40%, at least about 50%, at least about 60%, or at least about 75%, or more.

Alternatively, the sub-lethal concentration of the known antibiotic may be determined by measuring the activity of the target proliferation-required gene product rather than by measuring growth inhibition.

In order to characterize test compounds of interest, cells are contacted with a panel of known antibiotics at a sub-lethal level and one or more concentrations of the test compound. As a control, the cells are contacted with the same concentrations of the test compound alone. The IC_{50} of the test compound in the presence and absence of the known antibiotic is determined. If the IC_{50} of the test compound is substantially different in the presence and absence of the known drug then the test compound is a good candidate for use as an antibiotic. As discussed above, once a candidate compound is identified using the above methods its structure may be optimized using standard techniques such as combinatorial chemistry.

Representative known antibiotics which may be used in each of the above methods are provided in Table IV below. However, it will be appreciated that other antibiotics may also be used.

TABLE IV

Antibiotics and Their Targets

ANTIBIOTIC	INHIBITS/TARGET	RESISTANT MUTANTS
Inhibitors of Transcription		
Rifamycin, Rifampicin Rifabutin Rifaximin	Inhibits initiation of transcription/ß-subunit RNA polymerase, rpoB	гроВ, сгр, суаА
Streptolydigin	Accelerates transcription chain termination/ß-subunit RNA polymerase	rpoB
Streptovaricin	an acyclic ansamycin, inhibits RNA polymerase	rpoB
Actinomycin D+EDTA	Intercalates between 2 successive G-C pairs, rpoB, inhibits RNA synthesis	pldA

ANTIBIOTIC	INHIBITS/TARGET	RESISTANT MUTANTS
Inhibitors of Nucleic Acid N	/Ietabolism	
Quinolones,	subunit gyrase and/or topoisomerase	
Nalidixic acid Oxolinic acid	IV, gyrA	gyrAorB, icd, sloB
Fluoroquinolones	subunit gyrase,gyrA and/or	gyrA
Ciprofloxacin,	topoisomerase IV (probable target in	norA (efflux in
Norfloxacin	Staph)	Staph) hipQ
Coumerins	Inhibits ATPase activity of \$\beta\$-subunit	mpQ
Novobiocin	gyrase, gyrB	gyrB, cysB, cysE, nov, ompA
Coumermycin	Inhibits ATPase activity of ß-subunit	gyrB, hisW
A lhiaidin	gyrase, gyrB	
Albicidin	DNA synthesis	tsx (nucleoside channel)
Metronidazole	Causes single-strand breaks in DNA	nar
Inhibitors of Metabolic Pat	hways	
Sulfonamides,	blocks synthesis of	folP, gpt, pabA,
Sulfanilamide	dihydrofolate,dihydro-pteroate synthesis, folP	pabB, pabC
Trimethoprim,	Inhibits dihydrofolate reductase, folA	folA, thyA
Showdomycin	Nucleoside analogue capable of	nupC, pnp
	alkylating sulfhydryl groups, inhibitor of thymidylate synthetase	
Thiolactomycin	type II fatty acid synthase inhibitor	emrB
		fadB, emrB due to gene dosage
Psicofuranine	Adenosine glycoside antibiotic, target is GMP synthetase	guaA,B
Triclosan	Inhibits fatty acid synthesis	fabI (envM)
Diazoborines Isoniazid,	heterocyclic, contain boron, inhibit fatty	fabI (envM)
Ethionamide	acid synthesis, enoyl-ACP reductase, fabI	
Inhibitors of Translation	•	
Phenylpropanoids	Binds to ribosomal peptidyl transfer	
Chloramphenicol,	center preventing peptide translocation/	rrn, cmlA, marA,
	binds to S6, L3, L6, L14, L16, L25,	ompF, ompR
	L26, L27, but preferentially to L16	
Tetracyclines, type II	Binding to 30S ribosomal subunit, "A" si	
polyketides	on 30S subunit, blocks peptide	ompF
Minocycline	elongation, strongest binding to S7	
Doxycycline Macrolides (type I	Rinding to 50 S ribesomel subunit 275	
polyketides)	Binding to 50 S ribosomal subunit, 23S rRNA, blocks peptide translocation,	
Erythromycin,	L15, L4, L12	rrn, rplC, rplD, rplV,
Carbomycin,		mac mac
Spiramycin etc		

ANTIBIOTIC	INHIBITS/TARGET	RESISTANT
		MUTANTS
Aminoglycosides	Irreversible binding to 30S ribosomal	
Streptomycin,	subunit, prevents translation or causes	rpsL, strC,M, ubiF
Neomycin	mistranslation of mRNA/16S rRNA	atpA-E, ecfB, $hemAC$,D,E,G, $topA$,
Spectinomycin		rpsC,D,E, rrn, spcB atpA-atpE, cpxA,
Kanamycin		ecfB, $hemA,B,L$, $topA$
Kasugamycin		ksgA,B,C,D, rplB,K, rpsI,N,M,R rplF, ubiF
Gentamicin,		cpxA
Amikacin		rpsL
Paromycin		'PSL
Lincosamides	Binding to 50 S ribosomal subunit,	
Lincomycin, Clindamycin	blocks peptide translocation	linB, rplN,O, rpsG
Streptogramins	2 components, Streptogramins A&B,	
Virginiamycin,	bind to the 50S ribosomal subunit	
Pristinamycin	blocking peptide translocation and	
Synercid: quinupristin /dalfopristin	peptide bond formation	
Fusidanes	Inhibition of elongation factor G (EF-G)	fusA .
Fusidic Acid	prevents peptide translocation	
Kirromycin (Mocimycin)	Inhibition of elongation factor TU (EF-	tufA,B
Pulvomycin	Tu), prevents peptide bond formation Binds to and inhibits EF-TU	
Thiopeptin	Sulfur-containing antibiotic, inhibits	rplE
Tiamulin	protein synthesis, EF-G Inhibits protein synthesis	walC walD
Negamycin	Inhibits termination process of protein	rplC, rplD prfB
regamyem	synthesis	рIJB
Oxazolidinones Linezolid Isoniazid	23S rRNA	_
>T!: 0	* 1 11 1	pdx
Nitrofurantoin	Inhibits protein synthesis, nitroreductases convert nitrofurantoin to highly reactive electrophilic intermediates which attack bacterial ribosomal proteins	nfnA,B
Pseudomonic Acids	non-specifically Inhibition of isoleucyl tRNA	ileS
Mupirocin (Bactroban)	synthetase-used for Staph, topical cream, nasal spray	nes
Indolmycin Viomycin	Inhibits tryptophanyl-tRNA synthetase	trpS rrmA (23S rRNA methyltransferase; mutant has slow growth rate, slow
		chain elongation rate, and viomycin resistance)

ANTIBIOTIC	INHIBITS/TARGET	RESISTANT MUTANTS
Thiopeptides	Binds to L11-23S RNA complex	
Thiostrepton	Inhibits GTP hydrolysis by EF-G	
Micrococcin	Stimulates GTP hydrolysis by EF-G	

Inhibitors of Cell Walls/Membranes

B-lactams	Inhibition of one or more cell wall	
Penicillin, Ampicillin	transpeptidases, endopeptidases, and	C D E
Methicillin, Cephalosporins,	glycosidases (PBPs), of the 12 PBPs only 2 are essential: mrdA (PBP2) and ftsI (pbpB, PBP3)	ampC, ampD, ampE, envZ, galU, hipA, hipQ, ompC, ompF, ompR, ptsI, rfa, tolD, tolE tonB
• • •	Binds to and inactivates PBP2 (mrdA)	alaS, argS, crp, cyaA,
Mecillinam (amdinocillin)	Inactivates PBP3 (ftsI)	envB, mrdA,B, mreB,C,D
Aztreonam (Furazlocillin) Bacilysin, Tetaine	Dipeptide, inhib glucosamine	dppA
Bachysm, Tetame	synthase	арря
Glycopeptides Vancomycin,	Inhib G+ cell wall syn, binds to	
n	terminal D-ala-D-ala of pentapeptide,	
Polypeptides Bacitracin	Prevents dephosphorylation and regeneration of lipid carrier	ufn
Cyclic lipopeptide	Disrupts multiple aspects of	rfa
Daptomycin,	membrane function, including	
	peptidoglycan synthesis, lipoteichoic	
	acid synthesis, and the bacterial	
	membrane potential	
Cyclic polypeptides	Surfactant action disrupts cell	pmrA
Polymixin,	membrane lipids, binds lipid A	
Fosfomycin,	mioety of LPS Analogue of P-enolpyruvate, inhibits	murA, crp, cyaA
rosiomycm,	1 st step in peptidoglycan synthesis -	glpT, hipA, ptsI,
	UDP-N-acetylglucosamine	uhpT
	enolpyruvyl transferase, murA. Also	1
	acts as Immunosuppressant	
Cycloserine	Prevents formation of D-ala dimer,	hipA, $cycA$
Alafosfalin	inhibits D-ala ligase, ddlA,B	non A tun
Alaiosiailii	phosphonodipeptide, cell wall synthesis inhibitor, potentiator of ß-	pepA, tpp
	lactams	
Inhibitors of Protein Processin	g/Transport	
Globomycin	Inhibits signal peptidase II (cleaves	lpp, dnaE
	11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	44-

prolipoproteins subsequent to lipid modification, *lspA*

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It will be appreciated that the above cell-based assays may be performed using a sub-lethal concentration of a known antibiotic which acts against the product of any of the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi, or portions thereof, or homologous nucleic acids. In this way, the level or activity of a target, such as any of the proliferation-required polypeptides from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi, or homologous polypeptides may be reduced.

EXAMPLE 12

Transfer of Exogenous Nucleic Acid Sequences to other Bacterial Species

The ability of an antisense molecule identified in a first organism to inhibit the proliferation of a second organism (thereby confirming that a gene in the second organism which is homologous to the gene from the first organism is required for proliferation of the second organism) was validated using antisense nucleic acids which inhibit the growth of *E. coli* which were identified using methods similar to those described above. Expression vectors which inhibited growth of *E. coli* upon induction of antisense RNA expression with IPTG were transformed directly into *Enterobacter cloacae*, *Klebsiella pneumonia* or *Salmonella typhimurium*. The transformed cells were then assayed for growth inhibition according to the method of Example 1. After growth in liquid culture, cells were plated at various serial dilutions and a score determined by calculating the log difference in growth for INDUCED vs. UNINDUCED antisense RNA expression as determined by the maximum 10 fold dilution at which a colony was observed. The results of these experiments are listed below in Table V. If there was no effect of antisense RNA expression in a microorganism, the clone is minus in Table V. In contrast, a positive in Table V means that at least 10 fold more cells were required to observe a colony on the induced plate than on the non-induced plate under the conditions used and in that microorganism.

TABLE V

Sensitivity of Other Microorganisms to Antisense Nucleic Acids That Inhibit Proliferation in E. coli

Mol. No.	S. typhimurium	E. cloacae	K. pneumoniae
EcXA001	+	+	-
EcXA004	+	-	-
EcXA005	+	4	+
EcXA006	_	-	-
EcXA007	-	+	
EcXA008	+	-	+
EcXA009	-		-
EcXA010	+	+	+
EcXA011	-	+	-

EcXA012 - +	Mol. No.	S. typhimurium	E. cloacae	K. pneumoniae
BCXA013 + + + + + + BCXA014 + + + + + + + + + + + + + + + + + + +				
BeXA014		+		+
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EcXA067 - + -				
	EcXA067			<u> </u>

Mol. No.	S. typhimurium	E. cloacae	K. pneumoniae
EcXA068	-	-	-
EcXA069	-	+	_
EcXA070	-		-
EcXA071	+	_	-
EcXA072	+	-	+
EcXA073	+	+	+
EcXA074	+	+	+
EcXA075	+		
EcXA076	-	+	<u>·</u>
EcXA077	+	1 - 1	
EcXA077	+	+	
	+		+
EcXA080		-	-
EcXA082	-	+	
EcXA083	-	-	-
EcXA084	-	+	-
EcXA086	-	-	-
EcXA087	-	-	-
EcXA088	-		<u> </u>
EcXA089	-	<u> </u>	-
EcXA090	-	•	-
EcXA091	-	<u> </u>	-
EcXA092	-	-	-
EcXA093	-		-
EcXA094	+	+	+
EcXA095	+	+	-
EcXA096	-	-	-
EcXA097	+	<u>-</u>	_
EcXA098	+		-
EcXA099		_	-
EcXA100	-	-	-
EcXA101	-	_	_
EcXA102	-		
EcXA103	-	+	-
EcXA104	+	+	+
EcXA106	+	+	-
EcXA107	-	_	-
EcXA108	-	-	_
EcXA109	-	-	-
EcXA110	+	+	-
EcXA111	-	-	-
EcXA112		+	-
EcXA113	+	+	+
EcXA114	-	+	
EcXA115	-	+	_
EcXA116	+	+	-
EcXA117	+	<u>-</u>	-
EcXA118	-	-	_
EcXA119	+-	+	-
EcXA120	<u> </u>		-
EcXA121		- ;	-
EcXA122	+	-	+
EcXA123	+	-	
EcXA123			
EcXA125	-	-	-
10777123	<u> </u>	<u> </u>	·

Mol. No.	S. typhimurium	E. cloacae	K. pneumoniae
EcXA126	S. typnanariam	E. Cibacae	- x. preumonue
EcXA127	+	+	-
EcXA128	-	-	-
EcXA129		+	
EcXA130	+	+	
EcXA132	-	-	-
EcXA133		-	-
EcXA136	-		
EcXA137	-	-	-
EcXA138	+	-	-
EcXA139	<u> </u>		
EcXA140	+		-
EcXA141	+		-
EcXA142		-	-
EcXA143	-	+	
EcXA144	+	+	-
EcXA145	<u> </u>	-	-
EcXA146	-		
EcXA147	-	 	-
EcXA148	-		-
EcXA149	+	+	+
EcXA150	-	-	-
EcXA150	+	-	
EcXA151		-	-
EcXA153	+ +		
EcXA153	-		-
EcXA155			ND
EcXA156		+	ND
EcXA150 EcXA157	-	-	-
EcXA158	 		-
EcXA158 EcXA159	+	-	-
EcXA160	+	-	
EcXA162		-	-
EcXA163	 	-	-
EcXA163	-		-
EcXA165		-	-
EcXA166			-
EcXA166	-	-	-
EcXA168			-
EcXA169	<u> </u>		-
EcXA171	-		-
EcXA171	-	-	-
EcXA172		-	<u> </u>
EcXA174	-	-	
EcXA174			
	-	<u> </u>	<u>-</u>
EcXA176	+	<u>-</u>	
EcXA178	-	<u> </u>	-
EcXA179	+		-
EcXA180		-	-
EcXA181	-	-	-
EcXA182	-	-	-
EcXA183	-	-	
EcXA184	-		-
EcXA185	<u> </u>		-

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Mol. No.	S. typhimurium	E. cloacae	K. pneumoniae
EcXA186	-	-	-
EcXA187	+	+	+
EcXA189	+	-	-
EcXA190	+	+	+
EcXA191	+	+	-
EcXA192	-	+	-

Thus, the ability of an antisense nucleic acid which inhibits the proliferation of Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, 5 Helicobacter pylori, or Salmonella typhi to inhibit the growth of other organims may be evaluated by transforming the antisense nucleic acid directly into species other than the organism from which they were obtained. In particular, the ability of the antisense nucleic acid to inhibit the growth of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also 10 called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, 15 Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella 20 typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis or any species falling within the genera of any of the above species, may be evaluated. In some embodiments of the present invention, the ability of the antisense nucleic acid 25 to inhibit the growth of an organism other than E. coli may be evaluated. In such embodiments, the antisense nucleic acids are inserted into expression vectors functional in the organisms in which the antisense nucleic acids are evaluated.

It will be appreciated that the above methods for evaluating the ability of an antisense nucleic acid to inhibit the proliferation of a heterologous organism may be performed using antisense nucleic acids complementary to any of the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae,

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Helicobacter pylori, or Salmonella typhi (including antisense nucleic acids complementary to SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, such as the antisense nucleic acids of SEQ ID NOs.: 8-3795) or portions thereof, antisense nucleic acids complementary to homologous coding nucleic acids or portions thereof, or homologous antisense nucleic acids.

Those skilled in the art will appreciate that a negative result in a heterologous cell or microorganism does not mean that that cell or microorganism is missing that gene nor does it mean that the gene is unessential. However, a positive result means that the heterologous cell or microorganism contains a homologous gene which is required for proliferation of that cell or microorganism. The homologous gene may be obtained using the methods described herein. Those cells that are inhibited by antisense may be used in cell-based assays as described herein for the identification and characterization of compounds in order to develop antibiotics effective in these cells or microorganisms. Those skilled in the art will appreciate that an antisense molecule which works in the microorganism from which it was obtained will not always work in a heterologous cell or microorganism.

EXAMPLE 12A

Transfer of Exogenous Nucleic Acid Sequences to other Bacterial Species Using the Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi Expression Vectors or Expression Vectors Functional in Bacterial Species other than Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa,

Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae,
Helicobacter pylori, or Salmonella typhi.

The antisense nucleic acids that inhibit the growth of Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi, or portions thereof, may also be evaluated for their ability to inhibit the growth of cells or microorganisms other than Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi. For example, the antisense nucleic acids that inhibit the growth of Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi may be evaluated for their ability to inhibit the growth of other organisms. In particular, the ability of the antisense nucleic acid to inhibit the growth of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr

(also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae,
 Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium,
 Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus

dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis or any species falling within the genera of any of the above species may be evaluated. In some embodiments of the present invention, the ability of the antisense nucleic acid to inhibit the growth of an organism other than E. coli may be evaluated.

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In such methods, expression vectors in which the expression of an antisense nucleic acid that inhibits the growth of Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi is under the control of an inducible promoter are introduced into the cells or microorganisms in which they are to be evaluated. In some embodiments, the antisense nucleic acids may be evaluated in cells or microorganisms which are closely related to Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli; Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typh. The ability of these antisense nucleic acids to inhibit the growth of the related cells or microorganisms in the presence of the inducer is then measured.

For example, thirty-nine antisense nucleic acids which inhibited the growth of Staphylococcus aureus were identified using methods such as those described herein and were inserted into an expression vector such that their expression was under the control of a xylose-inducible Xyl-T5 promoter. A vector with Green Fluorescent Protein (GFP) under control of the Xyl-T5 promoter was used to show that expression from the Xyl-T5 promoter in Staphylococcus epidermidis was comparable to that in Staphylococcus aureus.

The vectors were introduced into *Staphylococcus epidermidis* by electroporation as follows: *Staphylococcus epidermidis* was grown in liquid culture to mid-log phase and then harvested by centrifugation. The cell pellet was resuspended in 1/3 culture volume of ice-cold EP buffer (0.625 M sucrose, 1 mM MgC1₂, pH=4.0), and then harvested again by centrifugation. The cell pellet was then resuspended with 1/40 volume EP buffer and allowed to incubate on ice for 1 hour. The cells

were then frozen for storage at -80°C. For electroporation, 50 µl of thawed electrocompetent cells were combined with 0.5 µg plasmid DNA and then subjected to an electrical pulse of 10 kV/cm, 25 uFarads, 200 ohm using a biorad gene pulser electroporation device. The cells were immediately resuspended with 200 µl outgrowth medium and incubated for 2 hours prior to plating on solid growth medium with drug selection to maintain the plasmid vector. Colonies resulting from overnight growth of these platings were selected, cultured in liquid medium with drug selection, and then subjected to dilution plating analysis as described for *Staphylococcus aureus* in Example 10 above to test growth sensitivity in the presence of the inducer xylose.

The results are shown in Table VI below. The first column indicates the Molecule Number of the *Staphylococcus aureus* antisense nucleic acid which was introduced into *Staphylococcus epidermidis*. The second column indicates whether the antisense nucleic acid inhibited the growth of *Staphylococcus epidermidis*, with a "+" indicating that growth was inhibited. Of the 39 *Staphylococcus aureus* antisense nucleic acids evaluated, 20 inhibited the growth of *Staphylococcus epidermidis*.

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TABLE VI
Sensitivity of Other Microorganisms to Antisense Nucleic Acids That Inhibit Proliferation of
Staphylococcus aureus

Mol. No.	S. epidermidis
SaXA005	+
SaXA007	+
SaXA008	+
SaXA009	+
SaXA010	+
SaXA011	-
SaXA012	-
SaXA013	-
SaXA015	+
SaXA017	-
SaXA022	+
SaXA023	-
SaXA024	-
SaXA025	+
SaXA026	+
SaXA027	-
SaXA027b	-

SaXA02c	-
SaXA028	-
SaXA029	+
SaXA030	+
SaXA032	+
SaXA033	+
SaXA034	-
SaXA035	+
SaXA037	+
SaXA039	-
SaXA042	-
SaXA043	-
SaXA044	-
SaXA045	+
SaXA051	+
SaXA053	-
SaXA056b	-
SaXA059a	+
SaXA060	-
SaXA061	+
SaXA062	+
SaXA063	-
SaXA065	-

Although the results shown above were obtained using a subset of the nucleic acids of the present invention, it will be appreciated that similar analyses may be performed using the other nucleic acids of the present invention to determine whether they inhibit the proliferation of cells or microorganisms other than Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi.

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Thus, it will be appreciated that the above methods for evaluating the ability of an antisense nucleic acid to inhibit the proliferation of a heterologous organism may be performed using antisense nucleic acids complementary to any of the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae,

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Helicobacter pylori, or Salmonella typhi, (including antisense nucleic acids complementary to SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, such as the antisense nucleic acids of SEQ ID NOs.: 8-3795) or portions thereof, antisense nucleic acids complementary to homologous coding nucleic acids or portions thereof, or homologous antisense nucleic acids.

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EXAMPLE 12C

As a demonstration of the methodology required to find homologues to an essential gene, nine prokaryotic organisms were analyzed and compared in detail. First, the most reliable source of gene sequences for each organism was assessed by conducting a survey of the public and private data sources. The nine organisms studied are *Escherichia coli*, *Haemophilus influenzae*, *Helicobacter pylori*, *Klebsiella pneumoniae*, *Pseudomonas aeruginosa*, *Staphylococcus aureus*, *Streptococcus pneumoniae* and *Salmonella typhi*. Full-length gene protein and nucleotide sequences for these organisms were assembled from various sources. For *Escherichia coli*, *Haemophilus influenzae* and *Helicobacter pylori*, gene sequences were adopted from the public sequencing projects, and derived from the GenPept 115 database (available from NCBI). For *Pseudomonas aeruginosa*, gene sequences were adopted from the Pseudomonas genome sequencing project (downloaded from http://www.pseudomonas.com). For *Klebsiella pneumoniae*, *Staphylococcus aureus*, *Streptococcus pneumoniae* and *Salmonella typhi*, genomic sequences from PathoSeq v 4.1 (Mar 2000 release) was reanalyzed for ORFs using the gene finding software GeneMark v 2.4a, which was purchased from GenePro Inc. 451 Bishop St., N.W., Suite B, Atlanta, GA, 30318, USA.

Subsequently, the essential genes found by the antisense methodology were compared to the derived proteomes of interest, in order to find all the homologous genes to a given gene. This comparison was done using the FASTA program v3.3. Genes were considered homologues if they were greater than 25% identical and the alignment between the two genes covered more than 70% of the length of one of the genes. The best homologue for each of the nine organisms, defined as the most significantly scoring match which also fulfilled the above criteria, was reported in Table VIIA. Table VIIA lists the best ORF identified as described above (column labelled LOCUSID), the SEQ ID, % identity, and the amount of the protein which aligns well with the query sequence (coverage) for the gene identified in each of the nine organisms evaluated as described above.

Table VIIB lists the PathoSeq cluster ID for genes identified as being required for proliferation in *Enterococcus faecalis, Escherichia coli, Pseudomonas aeruginosa*, and *Staphylococcus aureus* using the methods described herein. As indicated in the column labelled PathoSeq cluster ID, these sequences share homology to one another and were consequently grouped within the same PathoSeq cluster. Thus, the methods described herein identified genes required for proliferation in several species which share homology.

aisnoot	Data	Escherichia	Enterococcus	Haemophilus	Helicobacter	Klebsiella	Pseudomonas	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella	Streptococcus	Salmonella
		coli	faecalis	influenzae	pylori	pneumoniae aeruginosa		aureus	pneumoniae	typhi
EFA100001	SeqID	10430		86601	11603	11739		12309	13524	14040
	IDENTITY	27%	100	28	28%	75%		52%	25%	78%
•	COVERAGE	%66	100%	101%	79%	77%		%86	%86	%86
EFA100023	SeqID		10505					12860	13392	
	DENTITY		100%					27%	39%	
	COVERAGE		100%					%56	101%	
EFA100065 SeqID	SeqID	10322	_	111177	11351		12018	12820	13186	13733
	DENTITY	46%	100	46%	44%		48%	29%	%59	48%
	COVERAGE	%96	100%	%56			%16	%16	%86	%96
EFA100151	SeqID	10128		11247	11340		Ţ	•	13362	
	IDENTITY	20%	100%	37%	46%	110	46%	54%	51%	_
	COVERAGE	%66	100%	100%	100%		100%	%66	100%	
EFA100157	SeqID		10673		11448			12352	13176	
	DENTITY		100%		39%			64%	74%	
	COVERAGE		100%		%86			%86	%66	
EFA100165	SeqID				11564					14078
	IDENTITY	31%	90	33%	8		32%	73%	27%	73%
	COVERAGE	97%	100%	%86	100%		%96	%06	%96	%16
EFA100190	SeqID			19011		i				13966
	DENTITY	54%) 10	57	55%	55%	54%	78%	80%	24%
	COVERAGE	%00 I	101%	100%	%66	%06	100%	101%	101%	101%
EFA100194	SeqID				11426					14096
	DENTITY	%09	100	62	62		%09	85%	86%	61%
	COVERAGE	8001	101%	100%	102%		100%	101%	92%	101%
EFA100200	SeqID			11193						13731
	DENTITY	36%	9	38%			40%	20%	26%	39%
	COVERAGE	85%	100%	87%			85%	85%	%88	82%
EFA100210	SeqID			11104	11439					13968
	IDENTITY	53%	9	23%	53%		24%	74%	93%	53%
	COVERAGE	95%	101%	95%	94%		95%	101%	94%	95%
EFA100211	SeqID				11438		i		13205	
	COVERAGE	4070	1010%	4070	2970		4570	03%0	03%	
	COLUMN	0/ /0					0//0	21.70		

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TOCTION	Data	Prohousobia	Lutonococon	Homombiles	" Holiochaston	Vlahaialla	Dendomona	C		
	Time of the second	coli	faecalis	influenzae	nencooucier	neosieta pneumoniae aeruginosa	r seudomonas aeruginosa	лесте или рамен постем плетирище плеточен плетолница и ветанитом прирупососсия плетогососия принимента. Сой faecalis influenzae pylori preumoniae aeruginosa aureus pneumoniae typhi	sirepiococcus pneumoniae	samonena
EFA100289	SeqD DENTITY	30%	10810				31%			
EFA100295	SeqID	10045	10517	111	11601		11937	1239	13616	136
	IDENTITY COVERAGE	43%	100%	41%	41%		45% 97%	44%	45%	43%
EFA100312	SeqID IDENTITY COVERAGE		10641 100% 100%					12178 33% 88%		
EFA100329	SeqID IDENTITY COVERAGE		10782 100% 100%							
EFA100394	SeqID	10465	!	11238	11563		119611	13003	13684	13853
	IDENTITY COVERAGE	43%	100%	43%	42% 101%		108%	%66 %99	72%	44%
EFA100397	SeqID	10027	773					12396	13	14074
	COVERAGE	31%	100%	29% 98%		_	29%	43% 91%	46%	31%
EFA100399	SeqID	10295	766	11196	44				13413	13739
	COVERAGE	%86	100%	%86 6%60	%66 %60	_	28% 101%	%66 %7/	%9/. 100%	93% 98%
EFA100426	SeqID	10224	-			11638				13957
	COVERAGE	%66	100% 101%			%66 667		42% 91%	41%	28%
EFA100478	SeqID		786	Г	11338				13184	
	IDENTITY COVERAGE		100%	29% 72%	31% 70%			44%	43%	
EFA100615	SeqID			11139			12028	12641	13331	
	IDENTITY COVERAGE		100% 100%	44%			47%	61% 100%	78%	
EFA100617	SeqID		1	11216	11391					13765
	COVERAGE	43% 95%	100%	43% 96%	44% 78%		51%	63%	69% 82%	44% 93%
EFA100641	SeqID IDENTITY	10205 28%	10793 100				11896 31%		13334 32%	
\neg	COVERAGE	79%	100%				74%	85%	%28	

1						- 1				
TOCUSID	Data	Escherichia	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella	Haemophilus	Helicobacter		Pseudomonas	Pseudomonas Staphylococcus Streptococcus Salmonella	Streptococcus	Salmonella
		cott	faecalis	influenzae		pneumoniae aeruginosa	T		oniae	typhi
EFA100642	SeqID		10792		11520				13367	
	IDENTITY		100%		46%		46%	73%	%69	
	COVERAGE		%		100%		101%	100%		
EFA100668	SeqID	10026	Г	11184	11613		12013	12891	13505	14073
	IDENTITY	78%	100	28%	29%		28%	29%	20%	27%
	COVERAGE	83%	100%	%9L	78%		%76	82%	%66	%56
EFA100689	SeqID		10717					12523	13698	
	IDENTITY		100%					33%	33%	
	COVERAGE		100%					100%	100%	
EFA100704	SedID				11415				13171	13964
	IDENTITY	78%	90	78	11		75%	806	78%	7
	COVERAGE	100%	100%	100%	101%		101%	100%	101%	100%
EFA100739	SeqID	10111				11651	876		13220	14010
	DENTITY	71%	<u>8</u>	69	63%		71%	84%		70%
	COVERAGE	83%	101%	83%	%98	81%	83%	81%	82%	87%
EFA100740	SeqID					11633	942		13219	13717
	IDENTITY	45%	<u>≘</u>	47%	30%	45%	48%	64%	%09	44%
	COVERAGE	6	100%	94%	93%	94%	85%	94%	93%	94%
EFA100741	SeqID	10339			11430				13218	14098
	IDENTITY	40%	2	37	34	-	39%	48%	%09	40%
	KAGE.	103%	100%	102%	101%		102%	101%	100%	103%
EFA100742		10340			11431	••				14099
		25%	100	22%	36%		46%	%62	%88	52%
	RAGE	%66	101%	%66	%76		%66	101%	101%	%66
EFA100748	SeqID	10287						12595		13868
	COVERAGE	41%	100%	39% 99%	29%	42%	44%	52%		41%
EFA100756	SeqID	10112	10575		11396		11875	12327	13343	14009
_	IDENTITY	49%	100%		43%	_	45%	64%	62%	47%
	COVERAGE	75%	102%		75%		81%	94%	94%	75%
EFA100757	SeqID	10155	10897							
	DENTITY	27%	9							
	COVERAGE	%c8	%00I							
EFA100783	SeqID	10035	10811	10986	11543	. –		12738		13914
	COVERAGE	104%					78%	100%	%66 %67	99%

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	Saimoneua typhi					14042	46%	14065	40%	13874	45	101%	14093	48%				14094	46%	14027	30%	14086	26%	%68		
	orreptococcus saim pneumoniae typhi	13416 50% 101%	13641 85%	%96	13439 58% 99%		78%	230	63%	594	63%	<u>8</u>	3224	76%					70%	19191	56% 92%	13251	%98	89%	13554	101%
7	Excentrina Linerocciccus fraemopriaus relicodaties fixeoxidinomia resudamonius sugriyocciccus surepiococcias satinomiaia coli faecalis influenzae pylori pneumoniae aeruginosa aureus pneumoniae lipphi				12236 48% 98%		72%	368	59%	556	55%	101%	232	%66 82%					71%	12289	49%	715	76%	85%	504 71%	
1	Ateosteta Freudomorus pneumoniae aeruginosa		11775 63%	%26		5179	46%	12111	40%	1809	46%	101%	5158	43%					45% 102%	11801	39%	11945	%19	85%	11957 34%	- 1
11.1.11	nteostetta pneumoniae							11758	40%					49% 79%						11668	42%				11646 37%	- 1
प्राप्त त्वत्या	neucovacier pylori		11550 56%	89%		11410	52%	1 -	34%	1 4	40%	%66		42% 98%				11583	35% 98%	11607	29%	11401	20%	88%	11575 35%	83%
77	ndemopnius influenzae		11153 61%	95%			46%	11018	40%	127	45%	101%		48%					46% 99%	11210	40%	10982	28%	85%		
	Emerococcus faecalis	10863 100% 101%	10818 100%		10546 100% 101%	10627	100% 100%	62501	100%	1491	100%	100%		100%	10906	%001	100%		100%	18901	100%	10875	2		10722	101%
	coli		10382 62%	95%		T	47%	10399	40%	10269	۰	20		48%				10334	46%	10221	42%	10260	29%	85%		
	nana	SeqID IDENTITY COVERAGE	SeqID IDENTITY	COVERAGE	SeqID IDENTITY COVERAGE	SeqID	IDENTITY COVERAGE	SeqID	IDENTITY COVERAGE	SeqID	IDĖNTITY	COVERAGE	SeqID	IDENTITY COVERAGE	SeqID	IDÉNTITY	COVERAGE	SeqID	COVERAGE	SeqID	IDENTITY	SeqID	IDENTITY	COVERAGE	SeqID IDENTITY	COVERAGE
- 1	TOTOT	EFA100795	EFA100798		EFA100811	EFA100870		EFA100914		EFA100919			EFA100955		EFA100970			EFA100978		EFA100991		EFA101022			EFA101060	

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TOCTION	Data	Frahamahia	Dutangagan	LI.	II-linekant		7	r., 11.		11.0
	Dana	Escnericma 1.	Escrerichia Emerococcus Indemopnius Inelicopacier Meosiella	паеторишя	пенсовастег		rseuaomonas	r seudomonas Staphytococcus Streptococcus Salmonetta	Streptococcus	Salmonella
		coti	faecalis	influenzae		niae		aureus	pneumoniae	typhi
EFA101086	SeqID	10315	10763	11215				12953	13662	13764
	IDENTITY	37%	100	37%	27%	38%	35%	57%	25%	36%
	COVERAGE	91%	100%	%68	%86	%	92%	%86		93%
EFA101120	SeqID	10017	10687	11219	11331					14012
	IDENTITY	30%	100%	31%	27%		762	79%	64%	29%
	COVERAGE	102%	100%	102%	74%		103%	%66	%86	103%
EFA101121	SeqID		98901						13600	
	COVERAGE		100%					38% 98%	50% 99%	
EFA101123	SeqID	10420	Γ	11131	11478		11820	12674	13265	13783
	IDENTITY	43%	100	39%	33%	43%	40%	%0,	70,	42%
	COVERAGE	%86	100%	61%	%16	20	%96	%66	100%	%86
EFA101141	SeqID	10436			11573				13246	14045
	IDENTITY	35%	100	40%	32%		40%	%09	%02	31%
	COVERAGE	94%	101%	%96	%56		95%	%86	101%	%96
EFA101150	SeqID	10174		i	11556				13385	13943
	IDENTITY	35%	≅	36	56		33%	45%	280	36%
	COVERAGE	00	100%	100%	102%		100%	100%	100%	
EFA101159	SeqID				11442			12235	13197	13974
	IDENTITY	25%	≅	22	48%		49%	28%	%68	23%
	COVERAGE	800	101%	100%	81%		101%	%66	%66	100%
EFA101160	SeqID			11098	26511					13973
	IDENTITY	43%	≌	43%	33%	_	45%	62%	74%	43%
	COVERAGE	82%	100%	%26	%96		%76	100%	100%	63%
EFA101161	SeqID			11099						13972
	COVERAGE	39%	100%	35%			37%	%69	%99 1030/	36%
EFA101162	SeqID	10356	10555	11100	11441	11679	11993	12249	13200	13971
	IDENTITY	28%	100%	28%	26%	%65	%	%8	%	58%
	COVERAGE	100%	100%	100%	100%	100%	%66			100%
EFA101163	SeqID	10355			11594				13201	
	IDENTITY	%99	<u>ĕ</u>	% 89	9		20%	84%	906	
	RAGE	-	101%	%66	ı		100%	101%	100%	
EFA101164	SeqID	10354 55%	10558	11102	11593		5173	12258	13202	13970
	ы	91%					85%		97%	91%
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TOCOSID	Data	Escherichia	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella	Haemophilus	Helicobacter		seudomonas	Pseudomonas Staphylococcus Streptococcus Salmonella	Streptococcus	Salmonella
10000	4	con	Jaecaus	injinenzae		pneumoniae aeruginosa	П		ылав	ndha
EFA101165	SedID	10353	10559	11103	11592					13969
	IDENTITY	29%	100	%09	52%		61%	78%	88	26%
	COVERAGE	%56	100%	%56	%66		%56	100%	100%	%56
EFA101169	SeqID	10133	10574	16011			12025	12516		13849
	DENTITY	27%	100	28%		-	79%	41%		27%
	COVERAGE	93%	100%	%26			94%	100%		93%
EFA101253	SeqID	10389	10852	11065	11551		11838	13072	13457	
	IDENTITY	43%	100%	42%	31%		39%	54%	%19	
	COVERAGE	82%	100%	%26	%96		%66	61%	%66	
EFA101257	SeqID	10124	117	10976	4		11914	28	13357	14037
	IDENTITY	40%	100%	39%	39		37%	39%	28%	38%
	COVERAGE	%66	100%	%66	101%		%16	%16		101%
EFA101258	SeqID	10127	10918	10973	5		11892	02	13358	13871
	IDENTITY	40%	90	40%	36%		36%	41%	%99	23
	COVERAGE	97%	101%	%96	95%		%96	92%	95%	95%
EFA101322	SeqID		10620					-	13328	
	IDENTITY		100%					%99	%59	
,	COVERAGE		100%					%98	%98	
EFA101339	SeqID		10743		11448			12326	13391	
	IDENTITY		100%		33%	_		46%	%09	
	COVERAGE		100%		%16			%86	%86	
EFA101340	SeqID		10745							
	IDENTITY		100%						-	
	COVERAGE		102%	j						
EFA101354	SeqID	10047	10648		11608		_			13913
	COVERAGE	33%	100%	33%	32%		34%	38%	36%	32%
EFA101370	SeqID		10738				0/107	13126	7001	7701
	IDENTITY		100%					31%		
	COVERAGE		101%					%86		
EFA101403	SeqID		10662					12941		
	IDENTITY COVERAGE		100%				_	34% 100%		
EFA101404	SeqID	10210	10663	11214	11554		11921	12135	13418	13925
	IDENTITY COVER AGE	29%	100%	28%	39%		27%	59%	64%	30%
	CO V LINFAUL	2270					100%	77%	97.66	97%

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TOCOSID	Data	escherichia i.	Escherichia Enterococcus Haemophius Heilcobacter Klebstella	наеторишя	неисовастег		r seudomonas .	Pseudomonas Staphylococcus Streptococcus Salmonella	Srreptococcus	Salmonetta
		coli	faecalis	zae		pneumoniae aeruginosa			oniae	typnı
EFA101409	SeqID	10350	10524		11437				13207	
	DENTITY	54%	100	28%	44%		53%	%18	87%	
	COVERAGE	83%	101%	%08	%98		%16	%16	91%	
EFA101410	SeqID	10349	10525	11107	11436		5169	12216	13208	14108
	IDENTITY	%29	100%	64%	63%		%99	%06	%06	62%
	COVERAGE	101%	101%	101%	100%		100%	101%	101%	102%
EFA101411	SeqID	10348	10526	11108			5168		13209	14107
	DENTITY	20%	100%	43%			46%	%99	71%	46%
	COVERAGE	%16	%101	%16			93%	%96	%66	%16
EFA101412	SeqID	10347	10527							14106
	IDENTITY	%09	20	59	529	9	28%	85%	839	જ
	COVERAGE	100%	101%	100%	%86	101%	%66	%76	100%	
EFA101414	SeqID	10345	1		11435					14104
	IDENTITY	46%	9	47%	45%		46%	79%	819	49
	COVERAGE	%66	101%	%66	%66		100%	101%	101%	101%
EFA101415	SeqID	10344	_		11434					14103
	DENTITY	47%	100	20%	399		46%	63%	749	47%
	COVERAGE	%86	101%	%86	100%		%86	101%	101%	%86
EFA101416	SeqID	10343	10530		11433					14102
	DENTITY	20%	<u>)</u>	48%	42%		25%	%89	82%	51%
	COVERAGE	97%	101%	62%	91%		94%	%66	101%	%86
EFA101417	SeqID				11432					14101
	IDENTITY	25%	100	99%	61%		25%	72%	85%	55
	COVERAGE	100%	101%	%56	84%		92%	95%	94%	100%
EFA101424	SeqID	10220	10784	11276		11765	950			13934
	DENTITY	44%	100%	38%		34%	36%	65%	79%	41%
EEA 101475	COVERAGE	10240	10705	11076		13%	18%	101%	99%	39%
C2410142	Seque Transfer	10240	1000/	112/3			>			13803
	COVERAGE	%66		%66 %00			%66 866	93%	100%	4/%
EFA101477	SeqID	10263	10861	10965	11562		11948	13066	13525	14089
	DENTITY	25%	100	20%	41%		46%	26%	72%	20%
	COVERAGE	81%	100%	95%	%16		%56	94%	91%	%16
EFA101536	SeqID	10281	10823							
	COVERAGE	30%	100%							

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rocmen	D.42	Professionia	Produce of the second	77	Traffic Lanton	Victorialla	Promotomore	Chamber Longer	Chemitago	Calustan
	nin/	coli	faecalis	influenzae	neucooucier pylori	pneumoniae aeruginosa	aeruginosa	Lacher tenta taner occieda ingluenzae pylori pneumoniae aeruginosa aureus preumoniae typhi coli faecalis influenzae pylori pneumoniae aeruginosa aureus preumoniae typhi	pneumoniae	sumoneuu typhi
EFA101540	SeqID	10041	10487	11149	11456		11941		13438	13907
	IDENTITY	51%	100	20%	20%	_	46%	73%	<i>1</i> 6%	51%
	COVERAGE	92%			%98		92%			
EFA101541	SeqID	10042	10488	11150	11620		11940	12742	13437	13908
	IDENTITY	41%	100%	45%	35%		44%	%69	44%	41%
	COVERAGE	100%	100%	%86	121%		101%	100%	116%	100%
EFA101583	SeqID		10593							
	IDENTITY COVERAGE		100%							
EFA101670	SeqID		10511							
	IDENTITY COVER A CE		100%							
EFA101687	SealD	10238	10789	178	11517		11829	12811	13673	13864
-	IDENTITY	%	%(%	40%		%	%2.5	%	45%
	COVERAGE	%16					%16			%16
EFA101685	SeqID		10791		11369				13368	
	DENTITY		100%		47%		51%	62%	%69	
	COVERAGE	}	800		92%		%86	%/6	%66	
EFA101686	SeqID	10237	940	666	11325					13956
	IDENTITY	39%	100%	37%	37%		36%	64%	63%	38%
	COVERAGE	9,	100%	966	%66		%66	%66	%66	%66
EFA101695	SeqID	10204	629	017	11479	715				13928
	IDENTITY	34%	<u>ĕ</u>	32%	34%	31%	350	51%	75%	'n
	COVERAGE	إڅ	100%	106%	%9 <i>L</i>	93%	101%	100%	%66	105%
EFA101736	SeqID		_	11024						13976
	DENTITY	33%	100%	29%			27%	35%	32%	28%
EFA101737	SealD	10218	0778	11023			11923	7070	13341	13774
	IDENTITY	٠,	~	37%			,o	3%	%	58%
	COVERAGE	%86			_		%	100%	103%	
EFA101753	SeqID	10134	10552	11211			-			13826
	[IDENTITY	36%	100	37%			36%	20%	20%	37%
	COVERAGE	91%		%68		,	%06	94%	%66	%16
EFA101765	SeqID		10587						13353	
	COVERAGE		100%			-		%86 %97	92%	

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	Streptococcus	pneumoniae
	Staphylococcus	aureus
	Pseudomonas	aeruginosa
	Klebsiella	pneumoniae
TABLE VIIA	Helicobacter	pylori
TA	Haemophilus	influenzae
	Enterococcus	faecalis
	Escherichia	coli
	ata	
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TOCOSID	Data	Escherichia coli	Enterococcus faecalis	Haemophilus influenzae	Helicobacter pylori	Klebsiella Pseudomona pneumoniae aeruginosa	Pseudomonas aeruginosa	Escherichia Emerococcus Haemophitus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Isalmonella coli faecalis influenzae pylori pneumoniae aeruginosa aureus pneumoniae typhi	Streptococcus Salme pneumoniae typhi	Salmonella typhi
EFA101790	SeaTD	10414	10803	П						13747
	DENTITY	42%	100%				39%	46%		41%
	COVERAGE	101%		101%			2	101%		101%
EFA101791	SeqID		10804					12359		
	IDENTITY		100%					37%		_
	COVERAGE		101%					*		
EFA101792	SeqID	10030		11188	11458				13333	14077
	DENTITY	31%	2	32%	27%		33%	34%	47%	319
	COVERAGE	%86	100%	%96	%86		96%	101%	100%	%86
EFA101795	SeqID			11159						13886
	DENTITY	34%	10		36%		37%	36%	47%	
	COVERAGE	92	101%	%86	%66		%86	%86	%66	%16
EFA101797		10330	10924	091			12063	127	13364	13885
	DENTITY	23%	100%	529	49%		25%	29%		23%
	COVERAGE	92	100%	%86	%86		%86	%86	%66	%86
EFA101799	SeqID	10048	10926)14	11339		11934	806	13366	13897
	IDENTITY	23%	<u></u>	25%	49%	_	25%	54%	%99	54%
	COVERAGE	97%	100%	%26	94%		%16	%26		%16
EFA101833	SeqID	i	10720		11335		12039	12340	13451	14072
	IDENTITY	31%	100		36%		35%	21%	26%	31%
	COVERAGE	%6L	100%		%76		%68	%26	91%	19%
EFA101868	SeqID		10829							
	IDENTITY		100%							
	COVERAGE		100%							
EFA101872	SeqID									13779
	IDENTITY COVERAGE	62%	100%	62%	38%	%6Z 24%	%56 %09	93%	92%	62% 86%
EFA101873	SeqID		10816				11796			
	IDENTITY COVERAGE		100%				36%			
EFA101892	SeqID	10454	10506	11048	11281		12005	2142	13190	14021
	IDENTITY	47%	100%	47%	41%		53%	46%	46%	47%
	COVERAGE	100%	101%	100%	%26		100%		100%	100%
EFA101924	SeqID IDENTITY		10891 100%		11532 36%			12331 65%	13463 65%	
	COVERAGE		100%		101%			100%	94%	

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TOCUSID	Data	Escherichia	Enterococcus	Haemophilus	Helicobacter.	Klebsiella	Pseudomonas	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella	Streptococcus	Salmonella
		coli	faecalis	influenzae	pylori	pneumoniae aeruginosa		aureus	pneumoniae typhi	typhi
EFA101925	SeqID		10893					12332		
	IDENTITY		100%					26%		
	COVERAGE		100%					%66		
EFA101963	SeqID	10034	10848	11148	98511				_	13901
	IDENTITY	48%	100%	47%	46%		47%	21%	%69	48%
	COVERAGE	105%	100%	105%	%66		108%	101%	100%	105%
EFA102006	SeqID		10580				11830	12804	13315	
	IDENTITY		100%				33%	42%	43%	
	COVERAGE		100%		Ť		84%	%66	%56	
EFA102022	SeqID	10313	10881	11224		11754	12051		13485	13767
	IDENTITY	23%	100	23%	51%	54%	22%	78%	78%	52%
	COVERAGE	88%	101%	88%	87%	89%	88%	86%	%68	%68
EFA102023	SeqID									13768
	IDENTITY	51%	100	20%	38%	50,	20%	63%	70%	
ļ	COVERAGE	%86	100%	%66	%66	84%	97%	%66	99%	%26
EFA102091	SeqID	10363		11060	11568					13965
	DENTITY	%09	100	919	8		%29	75%	86%	55
	COVERAGE	101%	100%		100%		101%	100%	100%	101%
EFA102110	SeqID	10193	10841	11255			12082			13752
	IDENTITY	32%	<u></u>	34%			34%		62%	32%
	COVERAGE	103%	100%	94%			100%		100%	%66
EFA102183	SeqID				11330					13920
	IDENTITY	55%	100	24%			54%	%19	78%	55%
	COVERAGE	84%	100%	86%	85%		86%	%86	100%	84%
EFA102185	SeqID				11421	.632				13858
	IDENTITY	27%	<u>8</u>	79%	79%	28%	29%	63%	73%	27%
	COVERAGE	93%	101%	%06	94%	93%	%16	%16	%96	93%
EFA102186	SeqID	10448			11579					13817
	IDENTITY	29%	100	29%	27%			23%	% 99	30%
	COVERAGE	828	101%	%06				101%	95%	%06
EFA102205	SeqID		10769	\$8601	11375				13375	13997
_	IDENTITY	46%	100	38%	26%				25%	37%
- 1	COVERAGE	71%	102%	82%					%96	104%
EFA102253	SeqID			11175	11320				_	13865
	IDENTITY COYER A GE	53%	100%	55%	48%		53%	%19	%08 80%	54%
	COVERMUE	10070					101%	100%	9,4%	90%

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LOCUSID Data	Data	Escherichia coli	Enterococcus	наеторише	Helicobacter	Kiebsiella	Seudomonas	Escherichia Litterococcus Itaemophilus Helicopacter Klebsiella - Pseudomonas Staphylococcus Streptococcus Salmonella ooli Gaootio influenza milovi malavi maamanise ammanise ammanise kundi	Streptococcus Salm	Salmonella
000001144	1	202		injuentue		preumonne der ugmosu	П		pnemnounae	Wpin
EFA102282	SeqLD		10/29						13424	
	COVERAGE		100%					40%	46%	
EFA102338	SeqID			11012	11488		11954	12940	13272	13705
	IDENTITY	36%	100%	38%	35%		39%	42%	20%	38%
	COVERAGE	%56	100%	%26	%98		%86	%66	%66	%66
EFA102350	SeqID		10632							
	IDENTITY		100%							
	COVERAGE		101%							
EFA102351	SeqID		10634						13406	
	IDENTITY		100%					33%	38%	
	COVERAGE		100%					%16	101%	
EFA102352	SeqID			11186			12011	12347	13409	14075
	IDENTITY	40%	100	39%	35%	40%	39%	21%	25%	
	COVERAGE	101%	100%	101%	101%	101%	101%	%66	100%	101%
EFA102353	SeqID			11187	11329		Г		13398	14076
	IDENTITY	32%	100	34%	28%	_	32%	20%	61%	31%
	COVERAGE	%66	100%	%66	83%		%86	%86	%66	%66
EFA102389	SeqID	10378		11094			11781	12126	13263	
	IDENTITY	41%	100	42%			40%	24%	52%	
	COVERAGE	%16	100%	83%			%86	82%		
EFA102453	SeqID			10995		11762			13502	152
	DENTITY		100%	5	33%	33%		54%	54%	29%
	COVERAGE		101%	101%	%88	105%		101%	101%	
EFA102501	SeqID				11410		11997		13187	
	IDENTITY	45%	100	4	4	_	44%	75%	%92	4
	COVERAGE	112%	100%	111%	114%		113%	93%	%96	112%
EFA102502	SeqID		Г		11410					14042
	IDENTITY	47%	100	46%	25%		46%	72%	78%	46%
	COVERAGE	14%	100%	117%		-	116%	%66	%86	114%
EFA102503	SeqID		10643		11446				13481	13947
	IDENTITY	45%	100		37%		43%	61%	%59	419
	COVERAGE	%	100%		101%		101%	%86	100%	82%
EFA102518	SeqID	_	10647			11681				13881
	DENTITY	33%	9			50%		34%	54	32%
	COVERAGE	105%	2001			0/I/		102%	100%	105%

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	almonella vphi	13729	56% 77%	13732	%9L	100%			14007	58%	%96				13898	47%	%16				14011	25%	%96	13859	52%	2/2/			13822	71%	%66	13978	20%
	Streptococcus Salmo preumoniae typhi	13356 1	82% 81%	13261	%	100%			13221	%	101%	13216	63%	100%		%09	108%	13668	25%	100%	ī				81%	13401	71%	%66	13425	%08	100%		/4%
	Escherichia Enterococcus Haemophius Helicobacter Klebsiella Preudomonas Staphylococcus Streptococcus Salmonella coli faecalis influenzae pylori pneumoniae laeruginosa laureus pneumoniae typhi		%69 77%	12238	2%				12220	2%			62%	102%		21%			25%	100%					76%	0000			12590	%89	%66		08%0
	Pseudomonas aeruginosa	1	59%	12016	75%	95%	5159	%89	11000	62%		5161	45%	%26		48%	%66				11807	31%	%96	12074	54%	11943	51%	100%				12040	21%
	Klebsiella Pseudomona pneumoniae aeruginosa								11688	%																							
ABLE VIIA	Helicobacter pylori	11471	49%	11788	%29		11428	71%	11427	58%					11305	42%	99%							11420	52%	11300	44%					11362	0270
71	Haemophilus influenzae	11241	59%	11240	%02		11117	63%	11110	%19	%16	11115	40%	93%	11086	47%	%66				10956	%09	%96		23%	11205	52%	100%	11054	26%	%66	11261	0.770
	Enterococcus faecalis	10602	100%	10603	100%		10538	100%	0/ 501	%		10532	100	102%		2	100%	10734	100%	100%	10909	100%	100%		100%	10556	100%	100%	10478	00	100%	10896	102/20
	Escherichia	10327	59%	10326	۰,	95%		63%	10337	%	%96	10341	45%	93%		47%	%26					26%	%96		51% 89%	10285	~	%	Г	72%	%66	10142	0.00°
	Data	SeqID	IDENTITY	Seath	IDENTITY	COVERAGE	SeqID	IDENTITY COVED A GE	Sealth	IDENTITY	COVERAGE	SeqID	DENTITY	COVERAGE	SeqID	DENTITY	COVERAGE		IDENTITY	COVERAGE	SeqID	DENTITY	COVERAGE	SeqID	IDENTILY COVERAGE	SeqID	DENTITY	COVERAGE	SeqID	IDENTITY	COVERAGE	SeqID	I DENTIL I
- 1	TOCKSID	EFA102541		FFA 102542			EFA102549		FFA 102551			EFA102554			EFA102655			EFA102656			EFA102698			EFA102728		EFA102736			EFA102764			EFA102774	

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77%	10908	11167		2000			ادو	typhi
	106	112	112		51% 75% 11882	1263(133	139
	94% 101%	61% 93%			63% 94%	70% 93%	81% 96%	59% 94%
10274 66%	10854 100%	11154 64%	11298 58%		11932 64%	13128 74%	13313 83%	13866 65%
- 1	99% 100%	100%			100%			
10191 54%	· 10878 100%	11005 53%	11347 51%		11815 52%	12816 64%	13492 65%	13754 53%
100	100% 100%	100%			%66			
10297	106	10964	11323		11783	13090	13664	13737
7.001	100%	32%	%06 80%		31%	%86 %0c	%66 % 7 5	78% 100%
10434	106	11039	11413		11999	12451	13517	
65%	100%	66%	%09 %09		62%	86%	%98	
10221	10681	11210	11607	11668	11801	12289	13191	14027
0	100	40%	29%	vo.	%	49%	26%	30%
	91% 100%	93%	%86	94%	91%	93%	92%	93%
10435	10613	11038	11412		8611		13397	14046
~4% %4%	0 100%	3.2%	26%		51%	73%	73%	53%
	0000	,,,,,	9770		- 1	07001	-	99%
10293	10850	11041	11482	11728	11793	12541 73%	133 <i>7</i> 7 69%	13741
	%001 %66			%66	%66			
10437	10615	11072	11572		5180		13247	14044
59%	100%	64%	54%		65%	64%	68%	59
10767	10863	1008/	11402		11047	9970	12415	14000
41%	100%	41%	40%		41%		74%	40%
	85% 101%				%08		%56	
10251 32%	10689	10969 32%	11370 37%		11955 33%	12600	13518	13703
93	93% 100%				%96			

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LOCTISTD Data	Data	Fechorichia	Enterococcus	Haemonbilus	Holicohacter	Kloheiolla	Demondopues	Ecohenichia Emenocome Haemonbilue Helicohortee Klohviella Peandomonas Sambulococcue Stuentococcue Salmonalla	Strontococmic	Salmonolla
		coli	faecalis	influenzae	milari	nneumoniae aeruginosa	remainosa	smoon duding	nnoumoniae turbi	turbi
EFA103210	SeaID	10071						-	13319	13945
	IDENTITY	%95	100%	63%	39%		%	%6	%92	21%
	COVERAGE	%16	101%	%86		-	%16		101%	
EFA103268	SeqID	10365	10479	11062	11409		5178	12445	13231	13967
	IDENTITY	%69	100%	70%	%89		20%	83%	93%	40%
	COVERAGE	100%		100%			%66	101%	101%	
EFA103295	SeqID	10319	10633	11140	11493		12029	12640	13320	13771
	IDENTITY	%99	100%	28%	28%		%0/	79%	%98	%09
	COVERAGE	77%	101%	85%	%58		77%	100%	%96	92%
EFA103348	SeqID		10873	10983	11402		11946			
	IDENTITY		100%	36%	%65		39%			
	COVERAGE		103%	82%	82%		82%			
EFA103365	SeqID	10360	10533	11096	11443	11643	5177	12224	13196	13975
	IDENTITY	21%	2	28%	23%	28%	%85	82%	82%	28%
	COVERAGE	100%	101%	100%	%16	100%	100%	%88	101%	100%
EFA103375	SeqID	10177	09901	11222	11296		5120	12628	13302	
	IDENTITY	20%	100	52%	%98		20%	%99	78%	-
	COVERAGE	85%	102%	82%	%16		94%	102%	102%	
EFA103504	SeqID	10320	10671	11141	11492		12030	12638		13766
	IDENTITY	42%	100	45%	41%		48%	93%	81%	41%
	COVERAGE	%26	101%	97%	%96		%26	%86	100%	100%
EFA103508	SeqID		10672						13321	
	IDENTITY		100%						30%	
	COVERAGE		100%						80%	
EFA103571	SeqID	10335	62801	11121	11425		11988	12578	13240	14095
	IDENTITY	45%	100	47	48%		47%	%19	%89	45%
	COVERAGE	102%		102%	103%		102%		100%	102%
EFA103786	SeqID		10806					12361		
	IDENTITY		100%					%65		
	COVERAGE		100%					94%		

TABLE VII/

LOCUSID Data	Data	Escherichia coli	Enterococcus faecalis	Haemophilus influenzae	Helicobacter pylori	Klebsiella pneumoniae	Klebsiella Pseudomonas. pneumoniae aeruginosa	Esoherichia Enterococcus Haemophilus Helicobacter Klebsiella Peeudomonas Staphylococcus Streptococcus Salmonella coli faecalis influenzae pylori pneumoniae aeruginosa aureus	Streptococcus preumoniae	Salmonella typhi
SAU100040 SeqID IDEN' COVE	SeqID IDENTITY COVERAGE							12533 100% 101%		
SAU100053 SeqID IDENT COVEI	SeqID IDENTITY COVERAGE	10366 32% 97%	10504 46% 100%	11075 30% 99%	11376 32% 81%	11723 33% 84%	11855 33% 81%	12143 100% 100%	13318 48% 100%	13814 32% 97%
SAU100056 SeqID IDENT COVE	SeqID IDENTITY COVERAGE		10930 39% 98%					12 <i>577</i> 100% 100%	13477 33% 100%	
SAU100059	SeqID IDENTITY COVERAGE	10213 28% 71%	10598 70% 97%	11161 26% 95%	11528 26% 95%	11750 27% 71%	12064 28% 96%	12652 100% 100%	13433 25% 95%	13929 28% 71%
SAU100062 SeqID IDENI COVE	SeqID IDENTITY COVERAGE	10430 27% 103%	10618 52% 96%	10998 29% 103%	11603 29% 77%	11739 31% 76%		12309 100% 100%	13294 53% 97%	14040 28% 102%
SAU100077 SeqID IDENT COVE	SeqID IDENTITY COVERAGE		10565 64% 102%					12520 100% 100%	13464 62% 102%	
SAU100112 SeqID IDEN1 COVE	SeqID IDENTITY COVERAGE	% 97%			11477 52% 100%	11702 53% 77%	12096 46% 100%	12634 100% 100%		13895 49% 97%
SAU100114	SeqID IDENTITY COVERAGE	10152 44% 98%	10515 51% 88%	11279 43% 98%	11302 45% 98%		11851 43% 98%	12535 100% 100%	13387 25% 102%	13824 43% 98%
SAU100118 SeqID IDENT COVE	SeqID IDENTITY COVERAGE		10903 41% 101%				11828 27% 100%	12125 100% 100%	13262 37% 101%	
SAU100123 SeqID IDENT COVEI	SeqID IDENTITY COVERAGE	10258 52% 98%	10628 43% 100%	53% 53% 97%	11489 47% 96%		5192 52% 98%	12526 100% 100%	13421 45% 82%	14088 52% 98%
SAU100131	SeqID IDENTITY COVERAGE	10466 35% 71%		33% 97%			11960 40% 70%	12517 100% 100%		13854 35% 71%

					1ABLE VIIA	VIIA				
LOCUSID Data	Data	Escherichia coli	Enterococcus faecalis	Haemophilus influenzae	Helicobacter pylori	Klebsiella Pseudomona pneumoniae aeruginosa	Pseudomonas .	Escherichia Enterococcus Haemophilus Helicobacter Klebstella Pseudomonas Staphylococcus Streptococcus Salmonella coli faecalis influenzae pylori pneumoniae aerusinosa aureus pneumoniae tophi	Streptococcus pneumoniae	Salmonella tvvhi
SAU100133 SeqID	SeqID	10311	493	Ī		11703				13769
	IDENTITY	34%	44%	34%	33%	30%	31%	100%	43%	34%
	COVERAGE	%62	%66	%08	78%	82	79%	100%	%66	79%
SAU100139	SeqID	10355	557	11101	11594		74		13201	
	IDENTITY COVERAGE	65%	84%	66% 81%	64%		63%	100%	86%	
SAU100140	SeqID	10354	558	11102	440		5173	2258	3202	13970
	IDÉNTITY	9	%99	54%	40,		48%	100%	63%	54%
	COVERAGE	93	916	93	94%		93%	101	916	93%
SAU100141		10353	559	103	11592		5172	١.	203	13969
	IDEN III Y COVERAGE	%96 %cc	1019	%8c	34% 96%		%96 %96	100	/4% 100%	%96 860
SAU100157		10364	081	190	108	629	11996		232	13966
	IDENTITY COVER A GE	%09 100%	78%	%09 100%	55%	% 65%	57%	. 5	77%	60%
00 100		3	10170	10070	07.66	8	10076	Ini	101%	101%
SAU100158		10363 60%	481 75%	090 59%	11568 63%		.858 59%	443 100%	13233 77%	13965 58%
	RAGE	%86	97%	98%	%26		.0	100%	%16	
SAU100162 SeqID	1	10069 10	630	11239	11382					14084
	COVERAGE	43%	49%	44%	3/%		45%	100%	46%	45%
SATT100175	SedID	10250	27.70	11012	200		11054	19487	13070	12705
	DENTITY	vo.	2%	38%			%	%0	42%	35%
	COVERAGE	%86					2	100%		%66
SAU100182	SeqID							12362		
	COVERAGE							101%		
SAU100186 SeqID	SeqID			Γ	11423		11939	2317		13909
	IDENTITY COVERAGE	46%	61%	44%	46%		45%	100%	54% 99%	45% 101%
SAU100198 SeqID	SeqID				11445			2120	13414	
	DENTITY			-	29%			100%	29%	
SAU100227			10765					12525		
	COVERAGE		30%					100%		
SAU100242 SeqID IDENT	SeqID IDENTITY COVERAGE	10097 65% 94%		11201 62% 96%			11836 65% 95%	12336 100% 100%		14056 65%
SAU100246 SeqID	SealD		10821						13490	74/0
2001010	arhac.	_	17001	_	_	_	<u>-</u>		13450	_

Salmonella	mdka					13907 51% 88%				13769 40% 94%	13919 28% 99%	13711 27% 90%		13791 54% 96%
Streptococcus	pneumoniae 38% 93%					13438 65% 08%	13517 82%	13168 51% 97%		13491 49% 101%	13252 29% 99%	13244 40% 92%	13293 43% 100%	13521 74% 91%
Scherichia Enterococcus Haemophilus Helicobacter Klebstella Pseudomonas Staphylococcus Streptococcus Salmonella	anreus 100% 101%	12363 100% 100%	12122 100% 100%	12256 100% 101%	12141 100% 100%	12314 100%	2451 100%	12452 100% 101%	12453 100% 102%	12397 100% 100%	12313 100% 100%	12312 100% 100%	%001 100%	2358 100% 100%
Pseudomonas	pneumoniae aeruginosa					51%	%66 %59 1 66611	12000 42% 98%	12001 31% 103%	11885 40% 92%				12087 53% 97%
Klebsiella	риеитопіае							,0			11685 28% 99%			11727 55% 6 82%
Helicobacter Kleb	pytori					11621 51% 98%	=	⊏						11326 53% 96%
Haemophilus	apzuamifui					11149 47% 93%	689 88%	11083 41% 102%	11082 34% 93%	10990 38% _. 94%	lā	10963 30% 86%		11136 53% 96%
Enterococcus	35% 35% 101%				10617 26% 104%	9		10624 58% 98%		10774 50% 99%	10725 10 32% 100%	10814 44% 86%	10757 46% 99%	10802 73% 96%
Escherichia	700		10469 37% 88%			10041 52% 88%	10434 10 67% 99%	10433 41% 99%	10432 25% 92%	10311 40% 94%	10392 10 28% 99%	10013 26% 90%		10419 54% 96%
Data	IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE		SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE
LOCUSID Data			SAU100265	SAU100266	SAU100272 SeqID IDENT COVE	SAU100275	SAU100300			SAU100305	SAU100307	SAU100308	SAU100313	SAU100315 SeqID IDEN'I COVE

					אווא מחסטד					
LOCUSID Data	Data	Escherichia coli	Enterococcus faecalis	Haemophilus influenzae	Helicobacter pylori	Klebsiella Pseudomon pneumoniae aeruginosa	Pseudomonas aeruginosa	Escherichia Enterococcus Haemophitus Helicobacter Klebsiella Pseudomonas Stophylococcus Streptococcus Salmonella coli Jaecalis influenzae pylori pneumoniae aeruginosa aureus pneumoniae typhi	Streptococcus pneumoniae	Salmonella typhi
SAU100323 SeqID	SeqID	10216	10855					12575		13933
	IDENTITY COVERAGE	32%	71%)0% 100%		34% 88%
SAU100347	7.			10961					132	
	IDENTITY COVERAGE		44%	30% 84%			30% 100%	100% 1009	42% 100%	
SAU100355			10683					į .	13300	
	IDENTITY COVERAGE		42% 93%					100%	31%	
SAU100359			10757						13293	
	IDENTITY COVERAGE		52%					100% 100%	43%	
SAU100381	-	ì	10674					12276		14031
	COVERAGE	28%	29%				33%	100%		28% 101%
SAU100389	SeqID	10473	10737		11374			12279	13344	
	COVERAGE	75%	95%		%			100%	71%	
SAU100401	SeqID	10090	706	10980		11641		12576		14053
	COVERAGE	31%	95% 30%	21% 95%		33% 95%		100%		31% 99%
SAU100412		10102	563	94	11360		ĺ		13468	
	IDENTITY COVERAGE	31%	42%	30%	33%		35%	100%	40%	
CATTIONALA	Cecil	10453	1001	0. VO	- 12		110/12	100.	12401	13077
340100414	seque IDENTITY COVERAGE	60%	80% 80%	61% 98%	00511 60% 99%		91%	100%	13401 76% 969	%96 %09 %09
SAU100432 SeqID		10436	10614	071	⊨		5181	12450	13246	4045
	IDÉNTITY COVERAGE	34%	%86 %09	33%	31%		39%	100%	55%	31%
SAU100433 SeqID		10437	515	11072	115		5180	9449	13247	14044
	IDENTITY COVERAGE	58%	64%	63%	57% 99%		28% 98%	100%	%66 %69	58% 98%
SAU100436 SeqID	SeqID		10569					154	13393	
	COVERAGE		100%					100	27% 100%	
SAU100443 SeqID	SeqID . IDENTITY	10272 40%	10	1081 39%			11930 38%	12333 100%	13515 45%	869 40%
CAT 1100444	COVERAGE	95%	10507	%96	11540		92%	100%	100%	95%
jaao madaa jaadin	(Seque		10085	TIOTE	11340	_	/0611	12392	13403	14041

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LOCUSID Data	Escherichia	Enterococcus	Haemophilus	Helicobacter Kleb.	VIIA Klebsiella	seudomonas	Lesherichia Enterococcus Haemonhius Helicobacter Klebsiella Pseudomonas Stanbolococcus Strentococcus Salmonella	Streptococcus	Salmonella
ı	coli	faecalis	influenzae	pylori	pneumoniae aeruginosa	neruginosa .	aureus	pneumoniae	typhi
SeqID	10358		l	11595				13198	13973
DENTITY COVERAGE	41% 92%	62%	39% 97%	40%		46%	100%	63%	41% 93%
SeqID IDENTITY COVERAGE		10928 50% 99%					12565 100% 100%	13651 49% 99%	9
SeqID IDENTITY COVERAGE							12503 100% 100%		
SeqID IDENTITY COVERAGE							12121 100% 100%		
SAU100595 SeqID IDENTITY	10051 47%	10832 66%		11464 42%		12109 50%	12547 100%	46%	13722 42%
COVERAGE SeqID IDENTITY COVERAGE	36% 36% 99%	59% 50% 99%	11067 31% 100%	89% 11624 41% 92%	11656 38% 89%	93% 12110 42% 95%	100% 12548 100% 100%	90% 13173 30% 106%	91% 13720 32% 95%
SeqID IDENTITY COVERAGE							12616 100% 100%		
SeqID IDENTITY COVERAGE	10032 30% 102%	10870 61% 96%	11190 29% 100%	11349 29% 98%		12008 34% 87%	12293 100% 100%	507 50% 96%	14079 28% 104%
SeqID IDENTITY COVERAGE							12294 100% 100%		
SeqID IDENTITY COVERAGE	10378 44% 91%	10904 54% 88%	11094 43% 93%			11781 46% 73%	0%	13589 49% 89%	
SeqID IDENTITY COVERAGE		10502 26% 91%					12295 100% 100%	13314 25% 91%	_
SeqID IDENTITY COVERAGE	10079 27% 92%	10589 42% 103%			11698 2 25% 89%	5107 29% 101%	100%	13644 35% 105%	13724 26% 103%
SeqID IDENTITY COVERAGE	10051 50% 95%	10570 48% 94%		11464 46% 97%		12109 49% 95%	12168 100% 100%	13174 42% 95%	14109 50% 96%
SAU100658 SeqID	10322	10813	11177	11351		12018	12388	13186	13733

1										
LOCUSID Data	Data	Escherichia	Enterococcus	Наеторише	Helicobacter	Klebstella	Pseudomonas	Estrericnia Enterococcus Hamophilus Helicobacier Klebstella Pseudomonas Siaphylococcus Streptococcus Salmonella	Streptococcus	Salmonella
	IDENTITY	7001	Jaecalis Fon	ttuenzae 400/	pylori 460,	pneumoniae aeruginosa		aureus	рпеитопае	typni 1007
	COVERAGE	100%	39%	49% 100%	46%		48%	100%	38% 49% 100% 1	45%
SAU100659	92	10045	923		11601		11937 123	90	919	13911
		47%	54%	45%	40%		46%	100%	26%	44%
027001110	COVEKAGE	27%	0/76		105%	7	76	101%	3	- 13
SAU1006/9	Seqiii	10503		10997	453	713 220/	750/	37/	67.5	13/5/
	ίij	%96 %75		%66 %10	32%	96	976	100%	4270 104	%96 %65
SAU100684	SeqID	10412			11486		097	12632		13749
	COVERAGE	%16			%66 %04		%66 PV 04-	100%		%26
SAU100685	7							12633		
	IDENTITY COVERAGE							100%		
SAU100689	2		10694					12323	13311	
	IDENTITY COVERAGE		55% 98%					100%	46%	
SAU100702	102		10655					12196	13671	
	DENTITY		46%					100%	41%	
0411100710	_ _		2170					100%	91%	-
SAU100/10	Seqil						11908	12340		
	COVERAGE						739	1019		
SAU100714	SeqID IDENTITY	10465	10675	11238	11563		11961 12	100%	13382 13	13853 48%
	RAGE	108%					108%	1039	101%	
SAU100731	SeqID	10071	10688	11019	11371		11850	109	13319	
	IDENTILY COVERAGE	%66 %79	%6/ 100%	67% 100%	40% 1019		63%	100%	76% 100%	60% 101%
SAU100733	SeqID	10415			ı	11636	384	12602		13746
	COVERAGE	95%			33% 92%	42%	42%	100%		39% 95%
SAU100734 SeqID	SeqID	10321	10573	11142	l		331	503		13734
	COVERAGE	%86		97%	%06 %/7		%86 93%	100%	31%	29% 101%
SAU100736 SeqID IDENTITY COVERAG	SeqID IDENTITY COVERAGE		10585 27% 97%					1391 100% 100°	13404 26%	
SAU100738 SeqID	SeqID IDENTITY	10188 10 48%	847 45%	1953 46%	600 42%	634	907 51%	624 100%	45%	13981 49%
_	COVERAGE	97%	%86		97%	94%	976	100%	%26	94.6

					THOMAS ATTA	5				
rocosm	Data	Escherichia coli	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella coli faecalis influenzae pylori pneumoniae	Haemophilus influenzae	Helicobacter pylori	Klebsiella pneumoniae	Klebsiella Pseudomonas oneumoniae aeruginosa	Pseudomonas Staphylococcus Streptococcus Salmonella e aeruginosa aureus pneumoniae typhi	Streptococcus pneumoniae	Salmonella typhi
SAU100741		10081	10591		11459		11776	12409		13714
	IDENTITY COVERAGE	65% 100%	50%		35% 82%		54% 100%	100%		66% 101%
SAU100745	-	10442	10	(4)	11607	11733	1		453	13847
	IDENTITY COVER A GE	34%	53%	35%	31%	35%	34%	100%	49%	35%
SAU100747		200	10			101/0	3	2597	997	
			32%					100%	31%	
	COVERAGE		74%					100%	73%	
SAU100751		10425	10866	õ		11747	927	12335	-	13788
	COVERAGE	%66 %79	64% 99%	%86 88%		%Z9 82%	66 % 7.9	100%	63% 99%	%19 861%
SAU100752 SeqID	SeqID	10140					916	12524		14022
	IDENTITY COVERAGE	31%					35%	100%		38%
SAU100767	_	10290					12094	6/		13875
	IDENTITY COVERAGE	43% 100%					42% 90%	100%		42% 100%
SAU100771	SeqID	10084					11821	45	13306	13710
	IDENTITY COVERAGE	30%					29%	100%	28%	26%
SAU100773			0758	11093	11336	11763	11928	12377	13250	
	IDÉNTITY	47%	70	41%	41%	46%	51%	100%	20%	
0 1 1 1 0 0 1 1 1 0 0 0 1 1 1 0 0 0 0 0	COVEKAGE	%4%	100%	98%	%96	94%	93%	%10I	%96	
SAU100776 SeqID	SeqID							12482		
	COVERAGE							001		
SAU100778 SeqID	SeqID	10083		10957			11970	12514		14062
	COVERAGE	%68		%68 86%			%8			4/% 89%
SAU100793 SeqID	SeqID							12188	13392	
	COVERAGE							100%	2/%	
SAU100794 SeqID	SeqID	10203		3				12189		
	IDENTITY COVERAGE	25% 101%						100%		
SAU100799								12682 100%		
0.411100000	COVERAGE							100%		
lsacionologi Isacionologi	Cubec	_	_			_		12345		14081

Salmonella	typhi	35% 70%	14080	%96 %0¢	13765	50%	13811 42%	101%							¥	26% 104%				39% 106%	14026	95%	13704	38%	13754	100%		
Streptococcus	pneumoniae				l	58% 95%	13349						13183	100%	13601	26%		Ó	472	39% 100	473	36%	13506	48% 99%	13492	%66		
<u>IABLE VIIA</u> Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella	aureus	100%	12343	100%	Ç.	100%	12403		12212	100%	12211 100%	%	12210	~		100%	12329	100%	12401		12402	1009	.648	100%	1553	100%	12483 100%	
Pseudomonas	pneumoniae aeruginosa			%96 %	8615	48% 92%	12093 12 42%	%86													12071		11956	44%	11815	%		
VIIA Klebsiella	рпеитопіае															_			11719	37% 106%								
TABLE VIIA Helicobacter Kleb	pylori				10511	45% 91%									11342	28%			11367	35% 103%	11548	%96	11406	28% 101%	11347			
Haemophilus	influenzae				11216	4/%	11058 42%								É	28% 101%					11254	956	010	41% 100%	11005			
Enterococcus	faecalis				Č	63% 94%	10741 58%						10794	100%	10921	28%			10776	48% 98%		94%	877	49% 99%	878			
Escherichia	coli		10070	31% 94%	10314	4/%	10376 42%	%16								26%			Г	37% 106%	10446	94%	10252	39%	10191	100%		
Data		IDENTITY COVERAGE	SeqID	COVERAGE	SeqID	COVERAGE	SeqID IDENTITY	COVERAGE	SeqID				SeqID IDENTITY		•	IDENTITY COVERAGE	1	IDENTITY COVERAGE		IDENTITY COVERAGE	SeqID	RAGE	, All	COVERAGE	SeqID	COVERAGE	SeqID IDENTITY	
LOCUSID Data			SAU100810		SAU100813		SAU100831		SAU100836		SAU100838		SAU100839		SAU100843		SAU100845		SAU100858		SAU100859		SAU100865 SeqID		SAU100866		SAU100879 SeqID IDENTITY	

Salmonella typhi	14072 32% 85%	13733 43% 98%	14032 52% 93%	13957 38% 98%	13920 50% 85%			13876 32% 75%	13896 43% 91%	14074 32% 101%	14055 39% 101%	14012 39% 80%
Streptococcus pneumoniae	13451 45% 99%	13330 52% 989		348 52% 102%	13342 70% 96%					13478 34% 98%		13498 27% 83%
ococcus 100%	12340 100% 100%	12374 100% 100%	12376 100% 100%	139. 100% 100%	12138 100% 100%	12277 100% 100%	12278 100% 100%	12394 100% 101%	395 100% 100%	396 100% 100%	12615 100% 100%	12505 100% 100%
Klebsiella Pseudomonas Staphy pneumoniae aeruginosa aureus	12039 36% 81%	12018 45% 98%	12095 12 53% 92%	11905 1 36% 104%	11774 48% 83%				11983 30% 90%	33% 33% 96%	11834 39% 102%	120 <i>57</i> 39% 83%
Klebsiella		***************************************							11756 40% 86'			
Helicobacter Kleb pylori pneu	11335 35% 97%	11351 40% 99%	11509 52% 96%	11357 36% 99%	11330 49% 83%				11530 28% 91%			11506 36% 79%
Haemophilus influenzae		111 <i>77</i> 42% 98%	11001 53% 94%	11213 38% % 93%	11057 50% 82%				11191 31% 87%		11271 36% 101%	11219 36% 79%
Enterococcus faecalis	10720 51% 95%	10750 54% 98%	754 67% 749	701 60% 839	10952 51% 96%			10887 34% 72%	772 48% 86%	10773 43% 96%		10687 26% 108%
Escherichia coli	10429 31% 81%	10322 10 43% 98%	10410 10 52% 93%	10224 38% 97%	10393 50% 85%			75%	6 91%	%101 %	10095 39% 101%	10017 37% 80%
Data COVERAGE	TTY	TTY	SeqID IDENTITY COVERAGE	rity Rage	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE						
LOCUSID Data	SAU100880	SAU100882 SeqID IDENT COVE	SAU100885	SAU100886 SeqID IDENT COVE	SAU100887	SAU100899		SAU100916	SAU100920	SAU100921	SAU100932 SeqID IDENT COVE	SAU100944 SeqID IDENI COVE

LUCUSID Data	Data	Escherichia coli	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella coli faecalis influenzae pylori pneumoniae	Haemophilus influenzae	Helicobacter pylori		Pseudomonas aeruginosa	Pseudomonas Staphylococcus Streptococcus Salmonella aeruginosa aureus pneumoniae typhi	Streptococcus pneumoniae	Salmonella typhi
SAU100952 SeqID IDEN COVE	SeqID IDENTITY COVERAGE		10717 33% 104%					12523 100% 100%	13312 31% 102%	
SAU100959	SeqID IDENTITY COVERAGE		10704 58% 99%					12485 100% 100%	13504 49% 101%	
SAU100961	SeqID IDENTITY COVERAGE	10320 42% 98%	10671 63% 99%	47% 47%	11312 40% 97%		12030 50% 98%	12638 100% 1019	13322 57%	13766 42% 99%
SAU100962	SeqID IDENTITY COVERAGE				11299 28% 80%			12639 100% 101%	13 <i>577</i> 26%	
SAU100963	SeqID IDENTITY COVERAGE	10319 60% 84%	10633 79% 96%	11140 59% 81%	11493 61% 81%		12029 63% 84%	12640 100% 101%	13320 81% 92%	13771 60% 88%
SAU100964 SeqID IDENT COVE	SeqID IDENTITY COVERAGE		10501 61% 101%	11139 45% 76%			12028 47% 77%	.641 100% 100%	13331 60% 101%	i
	SeqID IDENTITY COVERAGE							12642 100% 101%		
SAU100970	SeqID IDENTITY COVERAGE	10128 52% 99%	10516 54% 99%	11247 39% 100%	11512 47% 100%		11891 52% 99%	12529 100% 100%	13362 46% 99%	
	SeqID IDENTITY COVERAGE		10686 38% 97%		11350 34% 73%			12606 100% 100%	13600 39%	
	SeqID IDENTITY COVERAGE	10185 29% 84%	10572 40% 98%	31% 31% 87%	 =		5122 26% 79%	12190 100% 100%		13820 30% 91%
SAU101020	SeqID IDENTITY COVERAGE							12710 100% 100%		
SAU101024	SeqID IDENTITY COVERAGE							12711 100% 101%		
SAU101028 SeqID IDENT COVE	SeqID IDENTITY COVERAGE	10034 11 46% 106%	10848 57% 101%	11148 43% 107%	11364 46% 100%		12006 46% 108%	12552 100% 100%	13471 55% 100%	13901 45% 106%
+cararave	OT bac		103/8	_	_	_	_	12608	13654	

Imonella	ohi			14027 31% 98%				13993 32% 88%				13827 37% 99%	13906 47% 99%	
ireptococcus Sa	pneumoniae typhi 37% 71%	13428 36%	N/COT	13191 14 46% 102%	13394 40% 99%	13380 32% 82%		1225 47% 101%	13666 49% 101%	13188 31% 97%		482 38% 96%	13	
<u>IABLE VIIA</u> Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus	aureus 100% 100%	12521 100%	12522 100%	12289 100% 100%	12290 100% 100%	30% 100%	12283 100% 100%	.84 100% 100%	.85 100% 100%	12191 100% 100%	~	100%	1968 12502 44% 100% 100% 100%	12299 100% 101%
Pseudomonas	pneumoniae aeruginosa	35%		1801 36% 98%			11974 35% 92%		34% 94%		827	11869 42% 99	11968 12502 44% 1009 100%	12070 43% 96%
V II.A Klebsiella	рпеитопіае			11668 38% 97%								11732 37% 99		
Helicobacter Kleb	pylori			11607 28% 108%				11462 37% 94%	11366 42% 74%			11404 37% 92%	11315 43% 98%	
Haemophilus	influenzae			11210 40% 100%			11156 34% 102%	11263 34% 88%				11248 39% 100%	11157 27% 82%	
Enterococcus	faecalis 36% 80%	10716 42%	ROS	10681 49% 103%	10682 41% 100%	10770 40% 89%				10755 36% 97%	10567 33% 96%	10768 45% 100%		10548 42% 98%
Escherichia	coli			10221 37% 98%			10066 36% 90%	10170 37% 89%			10450 35% 71%	10135 38% 98%	10040 47% 99%	
Data	IDENTITY		SeqD DENTITY	SeqD DENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE								
LOCUSID Data		SAU101038	SAU101039	SAU101065 SeqID IDENT	SAU101067	SAU101070	SAU101084	SAU101085	SAU101086			SAU101104	SAU101143	SAU101145 SeqID IDENT COVE

LOCUSID Data	Data	Escherichia coli	Enterococcus faecalis	Haemophilus influenzae	Helicobacter pylori	Klebsiella Pseudomonc pneumoniae aeruginosa	Pseudomonas aeruginosa	Escherichia Enterococcus Haemophitus Helicobacter Klebsiella Pseudomonas Siaphytococcus Streptococcus Islamonella coli faecalis influenzae pylori pneumoniae laeruginosa aureus pneumoniae typhi	Streptococcus pneumoniae	Salmonella typhi
SAU101155 SeqID	SeqID	10287		0	11352	11690	1	12310	1	13868
	IDÉNTITY COVERAGE	43%	49%	40%	30%	42%	42%	100%	37%	43%
041110116	COVERAGE	10/06	97.50	500	1	27.78	74.70	Ì		0000
SAUTOTION SEGIE	Seqiid	<u> </u>	%	60%	11333		%	100%		92/51
	COVERAGE	%%6			%16		%	101%		%96
SAU101159	SeqID IDENTITY		10891 65%		11532 36%			12331 100%	13463 54%	
			100%		100%			100%		
SAU101175								12213		
	IDENTITY COVERAGE							100%		
SAU101180 SeqID	SeqID		10888					12656		
	IDENTITY COVERAGE	38%	%68 86%				37%	100%		
SAU101183 SeqID	SeqID		10843					12304		
	COVERAGE		42% 102%					100%		
SAU101184 SeqID	SeqID	10477	10711	1218	11376	11735	12033	12305	466	13709
	IDENTITY COVERAGE	37%	46%	36% 102%	30%	38%	35%	100%	44%	38%
SAU101189								12264		
	IDĖNTITY COVERAGE							100%		
SAU101197				11024			1	12300		13976
	COVERAGE	98%	44% 98%	31% 101%			27% 100%	100%	46% 98%	30% 98%
SAU101198 SeqID	SeqID	10218	<u></u>	11023			11923	01	lco.	
	IDENTITY COVERAGE	43% 74%	20% 98%	43% 73%			41%	100%	46% 1029	
SAU101199 SeqID	SeqID	10088	<u> </u>	10970			11949	02	178	14052
	DENTITY	29%	40%	31%			36%	100%	37%	30%
SAU101220 SeqID	SeqID	10286	864					45	390	13870
	COVERAGE	32% 74%	37%					100%	39%	31% 74%
SAU101224					11533 28%			12 647 100%		
SATTIO1226 SeaTD	COVEKAGE		10837		%//	11658	11825	100%	13296	13701
) cody	_		_	_					17/61

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35%

57% 101%

3954

13299

12604

11845

11708

11521

18601

10684

|10232 | 35% | 95%

COVERAGE

IDÉNTITY

SeqID

SAU101275

100%

100%

102%

36%

46%

37% 100%

COVERAGE

DENTITY

SegID

SAU101271

11221

10719

(11556 25%

34% 94%

34% 96%

%86

33%

38% 93%

101%

101%

100%

46%

13385

112366 100%

11880

35%

27% 105% Escherichia Enterococcus (Haemophilus) Helicobacter Klebsiella | Pseudomonas Staphylococcus | Streptococcus | Salmonella 31% 47% 28% 20% 13864 4095 typhi 3837 101% 77% 97% 103% 85% 100% 98% %86 1<u>13359</u> 33% (<u>13238</u> 67% 1 oneumoniae 27% 35% 55% 21% 61% 35% 13486 13240 13383 13474 13317 38 47% 104% 12512 100% 12561 100% 100% 12573 100% 101% 100% 12570 100% 100% 12365 100% 100% 100% 100% 100% %001 100% 112564 100% 11<u>2488</u> 100% 1<u>12578</u> 100% 100% 100% 100% 12490 aureus 12303 36% 36% 33% 97% 52% 97% %68 100% %06 73% pneumoniae aeruginosa 32% 37% 43% 39% 12079 11829 11988 11881 1951 28% 75% 108% 29% TABLE VIIA 101% %86 105% 93% %86 33% 48% 47% 41% 45% pylori 11425 11399 11361 11324 11517 104% 100% %% 97% 27% influenzae 47% 47% 46% 11178 1108711121 11220 61% 100% 91% 84% 101% %66 %96 700 77% %66 10735 (<u>10616</u> 37% %19 32% 62% 52% 55% 57% faecalis 10513 10879 10789 10718 00501 10919 10175 50% 96% (10137 28% 73% 32% 101% 42% 48% 45% 100% 0238 10335 10089 0301 coli COVERAGE SeqID IDENTITY DÉNTITY DENTITY SedD SedID SedID SealD **GlpaS** Sea SealD SAU101266 SeqID LOCUSID Data SAU101236 SAU101239 SAU101270 SAU101235 SAU101240 SAU101242 SAU101262 SAU101267 SAU101247 SAU101231

171

					IABLE VIIA	<u>VIIA</u>				
LOCUSID Data	Data	Escherichia coli	Enterococcus faecalis	Haemophilus influenzae	Helicobacter . pylori	Klebsiella Pseudomom pneumoniae aeruginosa	Pseudomonas aeruginosa	Escherichia Enterococcus Haemophitus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella coli facalis influenzae pylori pneumoniae aeruginosa aureus preumoniae typhi	Streptococcus is pneumoniae	Salmonella typhi
SAU101286 SeqID	SeqID		10884					12292	13189	
	COVERAGE		4/% 100%					100%	40% 99%	
SAU101293	SeqID							12631		
	COVERAGE							101%		
SAU101300	SeqID		10751						13194	
	IDENTITY COVERAGE		57%					100%	54%	
SAU101301	SeqID		10752						13195	
	IDENTITY COVERAGE		%96 %/¢				27%			
SAU101302	SeqID		10753		11317			12559	13611	
	COVERAGE		49%		33%			~	72%	
SAU101310	SeqID		10924	11160	11321					13885
	IDENTITY COVERAGE	47%	52% 98%	48%	43%		47%	100%	51%	47%
SAU101311	SeqID	10094		11278			-	12563		13891
	IDENTITY COVERAGE	46%		46%			42%	100%		46% 95%
SAU101320	SeqID	10263	10861	10965	11562			12128	13254	14089
	IDENTITY	vo.	29%	49%			vo.	100%	26%	9
_	COVERAGE	%001	%66	%66	100%		%66	100%	976	100%
SAU101327	SeqID IDENTITY	10018 35%	10710 46%	11147			34%	512 100%	13495	14014 35%
	COVERAGE	%		101%			%76	101%	66	100%
SAU101339 SeqID	SeqID		10520		11365		11839	669		
	LUENTITY COVERAGE	%66 866	30%		26%		24% 97%	100%	%1.7 %	45% 99%
SAU101340	SeqID	10092					11840	14		13889
	COVERAGE	3/%					35%	100%		39% 104%
SAU101341	SeqID IDENTITY	10230	925	11212	11385		11898	100%	13365	13952
	COVERAGE	93%	%26	92%			%	100%	100%	
SAU101343 SeqID	SeqID IDENTITY	10422 50%	10649 55%	11162 49%		11721 50%			13346 58%	13785 51%
SAU101344 SeqID	SeaID	10171	10650	11252		99%	11826	12620	13347	13755
			_		_	-			_	-

Salmonella	38%	13894 36% 99%	13839 62% 100%	l i-i		13838 56% 98%	13874 45% 101%	13843 48% 99%				13862 52% 98%	13761 39% 94%	14067 32% 99%
Streptococcus	44%		13259 30% 91%	13286 55%	13285 59% 96%	13175 13 71% 13 % 101%	13295 50% 100%	179 569		13243 34% 77%	13432 41% 99%	13657 63% 96%	13422 37% 112%	13508 38% 92%
Escherichia Enterococcus Haemophilus Helicobacter Klebxiella Pseudomonas Staphylococcus Streptococcus Salmonella od Indendite Indiagnae	100%	12621 100% 100%	12622 100% 100%	12487 100% 100%	12486 13 100% 100%	2555 100% 1009	12556 100% 1009	12266 100% 100%	12274 100% 100%	12275 100% 100%	12145 100% 100%	12146 100% 100%	12147 100% 100%	12385 100% 100%
Pseudomonas	37%	11803 43% 99%	62% 100%	12069 46% 100%		11878 58% 98%	11809 45% 101%			11902 32% 79%		53% 53%	12055 38% 94%	12115 29% 98%
Klebsiella nneumoniae				٥		11684 55% 88%				.00		11635 39% 6 79%		32% 96%
Helicobacter Kleb		11282 35% 103%	11283 62% 101%	11318 32% 81%		35% 35% 97%	11577 40% 99%			11372 40% 86%		11292 42% 97%	11418 26% 98%	11368 27% 89%
Haemophilus	40% 79%		11163 29% 96%			10 <i>977</i> 54% 98%	11127 44% 101%					11179 50% 97%	11226 36% 97%	11030 319
Enterococcus	62%			10508 56% 98%	10507 60% 96%	10571 70% 101%	10491 55% 101%	10654 73% 98%				10707 60% 99%	10625 39% 90%	10830 52% 90%
Escherichia	48%	10058 36% 99%	10139 63% 100%	10184 61% 95%		10138 10 56% 88%	10269 45% 101%	10147 49% 99%			%86 98%	10239 10 53% 98%	10317 37% 102%	10403 10 33% 99%
Data	IDENTITY COVERAGE	SeqID IDENTITY COVERAGE												
LOCUSD Data			SAU101347	SAU101350			SAU101365	/	_					SAU101385

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		5			IABLE VIIA	ury				
LOCUSID Data	Data	Escherichia coli	Enterococcus faecalis	Haemophilus influenzae	Helicobacter pylori	Klebsiella Pseudomone pneumoniae aeruginosa	Pseudomonas, aeruginosa	Escherichia Enterococcus Haemophitus Helicobacter Klebsiella Preudomonas Staphylococcus Streptococcus Salmonella coli faecalis influenzae pylori pneumoniae aeruginosa aureus pneumoniae typhi	Streptococcus pneumoniae	Salmonella typhi
SAU101387 SeqID	SeqID	10402	10839		11549				i i	14068
	IDENTITY COVERAGE	27%	27% 35% 87% 88%		27%		27%	100%	32% 909	27%
SAU101389		10401	$\overline{}$	11029	11400		12113	187	13510	14069
	IDENTITY COVER A CE	55%	72%	57%	%09		57%	100%	74%	55%
SATT101398 SeoID	TANCE:	10313	881	774	202	754	051	100	747	191
accioiou.	ITY	55%	78%	54%	51%	57%	26%	100%	%89	ζŴ
	RAGE	100%	101	100	999	101	100	101%	101%	101%
SAU101399 SeqID		10312	882	686	116	755	050	325	669	15.0
	IDENTITY COVERAGE	%05 86%	63%	48% 98%	38%	51% 859	51% 97	100%	28% 99%	49%
SAU101400 SeqID	SeqID							326		
	DENTITY		46%		32%			100%	41%	•
	COVERAGE	-5			92%			%00I	76%	
SAU101408 SeqID IDENT	SeqID IDENTITY COVERAGE	10267 37% 100%	10509 43% 99%					12308	13278 42%	14050 39% 100%
101101110		2001	76701					1	IOI	100.0
	DENTITY COVERAGE		38% 38% 93%					100%		
SAU101427	SeqID							Г	13234	
	DENTITY							%0	48%	
	COVERAGE							100%	100%	
SAU101432				11046 57%	11286 60%	11744 63%	89% 89%	12184 100%	13538 26%	
	COVEKAGE			%66	100%	%I0I	99%	101%	73%	
SAU101436 SeqID IDENT	SeqID IDENTITY	10271 27%		11045 62%	1285 619		2067 59%	12183 100%		13873
	COVERAGE	%06		%66			8			%06
SAU101438 SeqID	SeqID	10146	825	11042					133	13842
	COVERAGE	30%	29% 949	29%		_		100%	27%	30%
SAU101444	SeqID	10254 10	827	11144	11301		12034	2381	13335	
	RAGE	100%	101%				%	100%	8	100%
SAU101445 SeqID IDENT	TTY	10248 52%		11207 52%			12037 54%	100%	13408 72%	13949 51%
SAU101446 SealD	SedID	10411	10674	2070			9	17383	100%	14031
			-	_	_			1	_	17071

DENTITY COVERAGE SeqID DENTITY COVERAGE SeqID DENTITY COVERAGE COVERAGE	COL		influenzae	miori	animonion	3	Sindano	opinomiona	huhi
	%86 %09	59% 100%	n)raenta	•	33% 37 97	%	100%	Direamonna	%66 80%
							12683 100% 101%		
	endiak A						12684 100% 100%		
SAU101455 SeqID IDENTITY COVERAGE							12686 100% 100%		
SAU101461 SeqID IDENTITY COVERAGE		10705 54% 93%			· · · · · · · · · · · · · · · · · · ·	11790 26% 86%	12680 100%		
rity Rage	10268 29% 77%	10708 45% 98%				11919 26% 91%	12679 100% 101%	584 26% 88%	14051 29% 77%
TTY	10469 10 38% 84%	905 29% 94%					12254 13 100% 100%	13454 25% 95%	13905 26% 73%
TTY	10125 109 40% 93%	39% 39% 95!	975 40% 96%	11290 32% 93%		11894 12 39% 96%	12130 100% 100%	41% 96	
ITTY	10126 10 55% 98%	51% 51% 1009	974 52% 98%	342 44% 98%	1738 36% 7	893 52% 9	11893 12123 13360 1409 52% 100% 48% 3 7% 98% 100% 99%	48%	14092 37% 101%
SAU101483 SeqID IDENTITY COVERAGE	10127 65% 88%	918 41% 909	973 59% 90%	11341 58% 90%		892 61% 8	12124 100% 101%	51% 929	13871 31% 94%
SAU101488 SeqID IDENTITY COVERAGE		10730 28% 95%				11868 25% 74%	11868 12164 1345 25% 100% 74% 100%	33% 98%	8 8
SeqID IDENTITY COVERAGE		104%					12165 100% 100%	5 42% 95%	
IITY RAGE	38% 38% 98%	10581 52% 101%	37% 37% 98%	11284 29% 78%	-	37% 37% 94%	12166 100% 101%	13323 43% 85%	13715 38% 98%
SAUI01493 [SeqID 11	10074 42%	-	41%	30%		11 832 43%	121 <i>67</i> 100%	13564 1 64%	13716 44%

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Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella	e pylori pneumoniae aeruginosa aureus pneumoniae typhi	% 101% 34% 101% 31% 31% 31%	11458 5187 12360 13333 14077 70% 32% 32% 32%	0% 86% 90% 100% 94%	12361	100%	100%	11712 12418 13249 36% 100% 49%	100%	1340	100% 34% 100% 89%		100%	13	38% 42% 100% 59% 46	100%	11867 12346 13406 140	27% 100% 32% 28%	75% 100.4% 100.4% 100.4%	27%	7% 83% 97% 100% 97%	234	100%	549 13460	38% 29% 100% 39%	70% 92% 102% 92%	11360	%86% 100% %88% %88%	11485 12551 13672 26% 160% 46%	70101
la Pseudom	niae aerugino)	5187						04%					12014	45%	1	11867	27%	10010	28%				12049	;	- 1	12019			
ter Klebsiel.	ounaud			%9				11712	=							%8					3%			11767		``		%6		10/
Helicobac	pylori		11458	3										11526	38%				0	27%							11360	3	11485	
Haemophilus	influenzae		11188 36%	2										11182	429		11183	27%	- 15	32%				11228	30%		76%	2		
Enterococcus	faecalis		10805 34%		10806	26%	100%			ğ	38%				63%	100%	ΙΦ.	33%	262	10030 50%	%66	10638	27%	762	38%	95%	10490		10485 48%	0207
Escherichia	coli	%96	10030	92%				10121 34%	104%					10024	41%	101%	10025	26%	10000	31%	<u>×</u>			10443	40%	%0/	10172	%16		
Data		COVERAGE	SeqID IDENTITY	COVERAGE	SeqID	IDENTITY	COVERAGE	SeqID IDENTITY	COVERAGE	SeqID	IDENTITY COVERAGE	SeqID	IDENTITY COVERAGE	SeqID	DENTITY	RAGE		IDENTITY	200	DENTITY	COVERAGE	SeqID	IDENTITY COVERAGE	SeqID	DÉNTITY	COVEKAGE	Seqilu	COVERAGE	SeqID IDENTITY	COVERAGE
LOCUSID Data		-	SAU101495 SeqID		SAU101497			SAU101509		SAU101526		SAU101529		SAU101541		_	SAU101543		0 411101545		_=	SAU101546 SeqID		SAU101549		-	Iccioines		SAU101554 SeqID	

						- 1		,		
TOCOSID IN	Data	Escherichia coli	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella coli faecalis influenzae pylori pneumonia	Haemophilus influenzae	Helicobacter pylori	67	Pseudomonas aeruginosa	Pseudomonas Staphylococcus Streptococcus Salmonella aeruginosa aureus pneumoniae typhi	Streptococcus pneumoniae	Salmonella typhi
SAU101561 S	SeqID	10400	160	Π	11355	11759	12112		307	14064
Ħ	IDENTITY	44%	57%	44%	38				49%	43
	COVERAGE	%66	%66	%66	100%	%66	100%	100%	%66	
SAU101565 S	SeqID	10134	55	11211			895	151	448	13826
<u> </u>	IDENTITY COVERAGE	3/%	%96 %05	35% 94%			36% 92%	100%	44% 99%	36% 92%
SAU101567 S	SeqID IDENTITY							12144		
۲	COVERAGE							100%		
SAU101570 SeqID	SeqID	10037	10690	11208		11700	11835	12584	563	13900
· ·	COVERAGE	%00I				34% 95%	33%	100%		30%
SAU101571 S	SeqID		9				11917	585	les	
<u> </u>	IDENTITY COVERAGE		45% 98%				33% 94%	100%	31%	
SAU101572 S	SeqID IDENTITY	10068	10692			11689	11864 12 43%	12586 13 100%	309	14083
	COVERAGE	75%	101%			%68	%96			75%
SAU101573 S	SeqID	10096	10693	Ċί			11865			14054
	IDENTITY COVERAGE	31%	49%	%86 %86			30% 101%	100%		31%
SAU101574 S	SeqID							12588		
	IDENTITY COVERAGE							100%		
SAU101575 S	SeqID		10869					12589	13638	
<u></u>	COVERAGE		91%					100%		
SAU101576 SeqID	eqID		10762					12554	4	
10	IDENTITY COVERAGE		32% 93%				79% 98%	100% 102%	39% 98%	
SAU101586 SeqID	keqID							12598	14	
10	COVERAGE							100%	34%	
SAU101592 SeqID	SeqID	10249	10605	10987	1555	11741	11952	12406	283	13950
	COVERAGE	101%		100%						101%
SAU101599 S II C	SeqID IDENTITY COVERAGE							12478 100% 100%		
SAU101610 SeqID	eqID	10449			11390		12048	12629		13816

LOCUSID Data	Data	Escherichia coli	Enterococcus faecalis	Haemophilus influenzae	Helicobacter Kleb	VIIA Klebsiella Pseudomond pneumoniae aerusinosa	Pseudomonas	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Isalmonella coli Italecalis Influenzae pylori Ineumoniae aerschosa Ianeus Inneumoniae Innei	Streptococcus	Salmonella
	IDENTITY COVERAGE	38%			% 101%		~	100%	and of the state o	38% 105%
SAU101612	SeqID IDENTITY COVERAGE							12637 100% 100%		
SAU101614	SeqID IDENTITY COVERAGE	10167 49% 100%	10678 55% 98%	11262 29% 93%	11534 29% 94%		39%	549 100%	13462 53% 99%	13851 48% 100%
SAU101616		10186 33% 102%	10667 28% 99%		11407 111 32% 88%	695 29% 1049	872 34% 96	432 100%		13903 33% 100%
SAU101622		10162 69% 100%			11619 29% 104%	710 67% 789	104 43% 101%	12430 100% 100%		13832 70% 100%
SAU101624	SeqID IDENTITY COVERAGE	10193 26% 101%		11255 27% 106%	11316 38% 97%				430 26% 103%	13752 26% 107%
SAU101630								12410 100% 100%		
SAU101632	SeqID IDENTITY COVERAGE							12407 100% 100%		
_	SeqID IDENTITY COVERAGE		10886 44% 99%					12201 100% 101%	13384 38% 98%	
	SeqID IDENTITY COVERAGE	10223 51% 92%					53% 53% 95%	193 100% 100%		
SAU101651	SeqID IDENTITY COVERAGE		10790 38% 97%		11552 28% 89%		12021 34% 90%	_	13369 42% 100%	
SAU101652	SeqID IDENTITY COVERAGE		10791 62% 97%		11369 49% 91%		12022 50% 95%	192 100% 100%	13368 56% 98%	
SAU101653	SeqID IDENTITY COVERAGE		10792 73% 100%		11520 46% 100%		12023 49% 100%		13367 63% 100%	
SAU101655 SeqID IDENT COVE	SeqID IDENTITY COVERAGE	31% 31% 84%	10793 50% 97%			-	30% 30% 83%	194 100% 100%	13334 33% 93%	

					TABLE VIIA	/IIA				
LOCUSID Data	Data	Escherichia coli	Enterococcus faecalis	Haemophilus influenzae	Helicobacter pylori	Klebsiella Pseudomon pneumoniae aeruginosa	Pseudomonas aeruginosa	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Preudomonas Staphylococcus Streptococcus Islamonella coli faecalis influenzae pylori pneumoniae aeruginosa aureus pneumoniae typhi	Streptococcus pneumoniae	Salmonella typhi
SAU101663	SeqID IDENTITY							12261		
								100%		
SAU101664	SeqID IDENTITY	10202	10512 41%	36%		-	11863	12262	13685	13823
		98%					%			%86 08%
SAU101674	94	10067						12594		14082
	IDENTITY COVERAGE	27%				·	27% 101%	100%		27%
SAU101679	93	10190	8	11055	11398		12105	12593	13264	13756
	IDENTITY COVERAGE	41%	53%	42%	36%		45%	100%	45%	40%
SAIT101681	SealD	10464	10746				11861	2502	13/10	13087
	IDENTITY	%					%	%0(%	40%
)	100%					2%	100%	102%	1
SAU101682	SeqID	10156	10670	11265						13884
	COVERAGE	~		102%				100%	34%	26% 94%
SAU101685	SeqID		10590				Γ	12152	13396	
	DENTITY		26%				37%	100%	26%	
C41101717	COVERAGE	00101	88%	2002	11010		y 7%	100%	3001	
	Sequi	>5	10380 51%	35%	31%		38%	12131	13352	140/0
	·	101%					%	100%	93%	
SAU101724	SeqID IDENTITY	10309	10588	11268	11337		12015	12136	13678	(L)
	COVERAGE	%					%(97%
SAU101726	(2)	10130	ğ	11026	11461		П	2134	13550	14
	IDENTITY COVERAGE	37% 101%	50%	42% 101%	36% 101%		40%	100%	48%	41%
SAU101727	SeqID		lō					12133	13551	
	IDENTITY COVERAGE		50% 101%					100%	49%	
SAU101728	SeqID IDENTITY	10019	10666	11053		11734	11800	12132	13182	14015
	COVERAGE	86%		88%		%	90%		74%	%98 86%
SAU101736 SeqID IDENT COVE	SeqID IDENTITY COVERAGE	10225 28% 72%					11817 38% 99%	12519 100% 100%		13958 29% 72%
SAU101737 SeqID	SeqID				11405		11817	12518		

	Salmonella			13706	31%	14043	46% 115%	14042	40% 116%	13967	65%	13934	41%	13863	48%							13900 44%	%16		14108 67% 102%
	Streptococcus Salm			165	45%	187	66 69%	646	101%	231	82%	280	%66 %86 %86	281	61% 101%	176	62% 98%	13308		13309	40%	13563 37%	%66	13207 79% 99%	13208 89% 101%
	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella coli facalis Influenzae mylori memmoniae pervorinsa aureus	100%	12367 100% 100%	12448	100%	7447	100%	446	100%	45	100%	50	100%	351	100%	2352	100%	12353	100%	12354	100	2355 100%	001	>2	216 100% 101%
	Klebsiella Pseudomonas	~				11997	45% 116%	5179	46%	5178	68%	950	36% 80	925	100			11917	93%	11916	41% 99%	866 42%	99%	70 55% 90	5169 66% 100%
VIIA	Klebsiella pneumoniae			11671	30%							765	35% 82%							11689	%68 89%	11700 11 35%	92%		
TABLE VIIA	Helicobacter nylori	32%					40% 120%	11571	%08 80%	11409	65%			11294	27% 77%	11448	43% 88%							.437 48% 86%	62% 100%
	Haemophilus influenzae					11037	47% 114%	11036	117%	11062	66%	11276	37%	11275	51% 100%							11208 45%	%/6	11106 55% 86%	11107 69% 101%
	Enterococcus faecalis		10562 44% 101%	ΙΦ	46%	10626	75% 99%	627	100	479	83%	784	65% 1019	785	63% 1019	10673	64% 97%	10495	%66	10496	100%	10498 11	100%	10524 81% % 89%	525 90% 101%
	Escherichia coli			10474	30%	10438	12%	10439	116%	10365	65%	10220	43%	10240	50% 100%							10037 110	7	10350 10 51% 86%	10349 67% 101%
	Data	IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID	IDENTITY COVERAGE	-	IDENTITY COVERAGE	SeqID	COVERAGE	SeqID	COVERAGE		IDENTITY COVERAGE	SeqID	IDENTITY COVERAGE	SeqID	IDENTITY COVERAGE	SeqID IDENTITY		SeqID			CUVEKAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE
	LOCUSID Data		SAU101744	SAU101751 SeqID		SAU101752		SAU101754 SeqID		SAU101756 SeqID		SAU101771		SAU101772		SAU101777 SeqID		SAU101781		SAU101782		SAU101784	000000000000000000000000000000000000000	SAU101790	SAU101791 SeqID IDENT COVE

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					TENDER VIII					
LOCUSID Data	Data	Escherichia coli	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella coli	Haemophilus influenzae	Helicobacter		Pseudomonas	Pseudomonas Staphylococcus Streptococcus Salmonella	Streptococcus	Salmonella
CAT1101700	Coom	10240	10506	T	Prote	pricamound	1	12217	Т	14107
SAU101/92	Sequi	10348	1020	11108				1000/	,	1410/
	COVERAGE	%96 6%	94%	%56 62%			45%	100%	94%	%0c
SA11101793	SeaTD	10347	1527	11109	11589	11654	5167	2218	13210	14106
	IDENTITY	64%	85%	%	%	vo	3%	%0	79%	64%
	田	8	101%	%66		101	%66	101	100%	
SAU101795	SeqID	10345	10528	11111	14		П	ı	13212	14104
	DENTITY	21%	79	47%	44%		44%	100%	%92	51%
	COVERAGE	8	101%	%66	%86		%	101	101%	101%
SAU101797	7.	10343	1530	113	7			12221	13214	14102
	IDENTITY COVERAGE	45%	68%	41%	41%		48%	100%	66%	46%
0.411101700	COLUMN	102.47	10170	114	435		2100	101	216	14101
94/101/30	DENTITY	>	77%	55%	11452		%	7777	66%	14101
	Ħ	%66	95	%66			%66	1015	%96	
SAU101799	SeqID	10341 10	532	11115			5161	2223	2	
	斑	100%	102	100%			%	100%	%86 %60	
SAU101800	SeqID	10340	534	116				12225		14099
		47%	79%	46%	9		_	100%	84%	47%
0000	COVERAGE	07.66	IOI	266	30%		0	101	101%	35%
SAU101802	Seqili	100/5	556	308 52%	348 31%	11633	11942	1000%	13219	13717
	ΙÜ	97%	97.	979	93.	97.4	84%	100	96	%26
SAU101803	SeqID	10111	537	352	429	1651	928		13220	14010
	IDENTITY	71%	84% 1019	71%	60% 1001	70%	71%	100%	82%	70%
SAU101805 SeaTD		10337	539	11119	427		066	2229	13221	14097
	TTY	53% 96%	75% 101%	52% 99%	58% 99%		%96 %09	100%	74%	52%
SAU101806	SeqID	10336 10	540	1120	17		П	12230	3222	14096
		%79	85%	64%	9		%19	100%	85%	63%
	COVERAGE	100%	101%	100%	102%		100%	101	92%	101%
SAU101807	SeqID	10334	541	1122	11583		11987	12231	13223	14094
	COVERAGE	%66 %7±		%66 60%			%66 874	පි	%66 80%	%66 60%
SAU101808		10333		1123	<u>ı∽</u>	11627	58			14
	RAGE	48% 98%	65% 103%	49% 98%	46% 99%	48% 78%	45% 98%	100%	67% 106%	48% 98%
SAU101810 SeqID		10053	544	1229	1625	11666	606	12233	13441	14110

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	s Salmonella	sypni 36%	73%	13721	34%	12720	13/29	%l %66 %l	13732	%66 %		~			13775	44%	13924	28%	% 94%	13999	%66 %	13953	20% 101%	13713	56% 104%			13797 39% % 98%	
	Streptococcu		%88	13440	45%	113356	65%	666	13361		13494	35%			135	46%	13291	32%	834	13445 13	82,8	544	102	1379	75%		.0	13305 62% 99%	
	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Preudomonas Staphylococcus Streptococcus Salmonella	aureus pn	100%	12234	32% 100%	1007	16237	101%	12238		12369	100%	'n	100%	12373	100%	105	100%	100	510 100%	1000	506	100%		100%	12569 100%	100%	12571 100% 100%	12572
	Pseudomonas	rugmosa 33%	72%	11888	32%	6100	55%	%16	12016	্		31%	12004	28%	11794	35%	12100	25%	86	855 47%	9	899	101%	12058	56%			11802 39% 98%	
VIIA	Klebsiella	pneumoniae ae	2	99	33%	9 9	26%													11723 111	946				_				
IABLE VIIA	Helicobacter	pyiori pne	77	163	32% 2	70	+/1 47%	32	288		11307	%55 80%			11481	44%				376 48%	66	567	102%	11472	56% 101%			11334 33% 101%	
	Haemophilus	mfluenzae py 34%	787	11068	33%	11041	57%	<u>\$</u>	11240	%86	11231	32% 95%			12	28%	11236	32%	8	075 33%	95%			2	54% 103%			10955 40% 98%	
	Enterococcus	faecalis 57%	88	545	49%	50	700 700	96							1	49%	849	33%	780	942 70%	95%	739	47%		77% 99%			10817 63% 100%	
	Escherichia	25% Jae	%9 <i>L</i>	10196	38%	10227	58%	8	10326	%86			10158	33%	10207	42%	10398	%	94%	10105 10	98%	10231	30%	10015	56% 103%			102 <i>57</i> 10 40% 98%	
		IDENTITY	田		IDENTITY COVED A CE	ا		COVERAGE	SeqID InFNTITY	COVERAGE	SeqID	IDENTILY COVERAGE	SeqID	IDENTITY COVERAGE	SeqID	IDENTITY COVERAGE			RAGE	, ALL	COVERAGE	SeqID	COVERAGE	SeqID	IDENTITY COVERAGE	SeqID IDENTITY		SeqID IDENTITY COVERAGE	SeqID IDENTITY
	LOCUSID Data			SAU101811		0 4 7 7 1 0 1 0 1 4	SACIOIS14 SEGID		SAU101815		SAU101818 SeqID		SAU101824 SeqID		SAU101833 SeqID		SA11101839			SAU101842 SeqID		SAU101845		SAU101849		SAU101857 SeqID		SAU101862	SAU101864 SeqID IDENT

					TABLE VIIA	/ <u>IIA</u>				
LOCUSID Data	Data	Escherichia coli	Enterococcus faecalis	Haemophilus influenzae	Helicobacter pylori	Klebsiella Pseudomon pneumoniae aeruginosa	Pseudomonas aeruginosa	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Peeudomonas Staphylococcus Streptococcus Sdinonella coli faecalis influenzae pylori pneumoniae aeruginosa aureus	Streptococcus pneumoniae	Salmonella typhi
SAU101865	SeqID	10044		Г			Τ	12318		13910
	IDENTITY	43%	28%	45%	40%		40%	100%	54%	41%
	COVERAGE	85%	%88	%88	%18		87%	100%	%88	88%
SAU101866 SeqID	SeqID		10835				11873	12319	13586	
- 1	COVERAGE		42%				%	100%		
SAU101868 SeqID	SeqID	10049	10733	<u></u>	11305					13898
	IDENTITY COVERAGE	45%	%95	45%	42%		48%	100%	49%	45%
SAT1101869	SeaTD		10734	0/101			2001	12321	13668	7///
	DENTITY		55%					100%	49%	****
	COVERAGE		%001					100%		
SAU101876	SeqID							12169		
·	IDENTITY COVERAGE							100%		
SAU101881	SeqID	10325						12162		13728
	IDENTITY COVERAGE	42%				•	41%	100%		42%
SAU101882	SeqID	10246	108					12163		13727
	IDENTITY	33%	30%			31%	31%	100%		
	COVERAGE	%96	%68			73%	94%	100%		%56
SAU101890	SeqID	10374		11125				12280		13809
	IDENTITY COVERAGE	23%		49%	1-11		47%	100%		53%
SAU101891	SedID	10295	9920	11196	11483		16/11	12281	13413	13739
	DENTITY		72%	%	%09		%	100%	%29	64%
	COVERAGE	12%	816	%06	%06		93%	100	92%	%16
SAU101893	SeqID	10300				11748	1981	282	3290	13825
	COVERAGE	40% 87%	4/%			41%	35% 93%	100%	40% 95%	43%
SAU101904	SeqID	10047	648		4		1935	617	345	IQ.
	DENTITY	34%	38%	33%	31%		31%	100%	34%	33%
CATTINION7	COVERAGE	10367	10170		115		11005	100%	93%	98%
	DENTITY	×°	70-407 60%		741		73%	100%	2%	74%
	COVERAGE	100%	101%	%00I			~	100%	101%	100%
SAU101909	SeqLD	10390		11249 32%	11346 29%		36%	12441		14063
0,000	COVERAGE	%66		88%			93%	100%		73%
SAU101910 SeqID	SeqID	10199	_	_	_	_	11818	12440		

Salmonella	ndka				14003 45% 88%	13998 31% 76%			13956 37% 98%	13708 48% 97%	14088 47% 105%			
Streptococcus	oneumoniae						13500 25% 80%	13386 51% 74%	13455 13956 58% 37% 0% 100% 98%	13241 64% 97%	13636 46% 98%			13260 47%
Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella	aureus 100% 100%	12439 100% 100%	12438 100% 100%	12709 100% 100%	12186 100% 101%	12187 100% 100%	12454 135 100% 101%	11957 12455 1338 57% 100% 76% 100%	11901 12456 134 35% 100% 134 99% 100%	12423 100% 100%	12424 100% 100%	12425 100% 100%	12426 100% 101%	12427 100%
Pseudomonas 2	~				1897 45% 88	30% 30% 83		11957 57% 76%	11901 35% 99%	12035 51% 96%	11787 12 49% 98%			
Klebsiella	pneumoniae aeruginosa 60% 97º				705 43% 88%			646 46%						
Helicobacter Kleb	pytori				538 37% 86%	11480 27% 88%		575 58% 72%	11325 38% 98%	11304 49% 91%	11489 43% 105%			11555 28%
Haemophilus	mjruenzae				11007 32% 92%	33% 33% 90%		56 49% 739	36% 36%	59 46% 98	11134 47% 106			11267 44%
Enterococcus	aecans	10838 26% 90%			31% 91%	10568 31% 4 92%	10938 40% 101%	10388 10939 1110 46% 47% 72% 78%	10237 10940 38% 64% 98% 99%	10941 61% 98%	58% 98°			
Escherichia	con 56% 97%				10101 45% 88%	10106 30% 90%		10388 46% 72%	10237 38% 98%	10476 109 48% 97%	10258 47% 105%			
Data	IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	TITY	ITTY RAGE	SeqID IDENTITY COVERAGE	TTY	TTY	SeqID IDENTITY COVERAGE	ш	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY
LOCUSID Data		SAU101915	SAU101922	SAU101948	SAU101966 SeqID IDENT COVE	SAU101968 SeqID IDENT COVE	SAU101991	SAU101995	SAU101996 SeqID IDENT COVE	SAU101999	SAU102001	SAU102002 SeqID IDENTITY COVERAG	SAU102003	SAU102006 SeqID

					1111	****				
LOCUSID Data	Data	Escherichia	Enterococcus footalis	Haemophilus influenzae	Helicobacter	Klebsiella	Klebsiella Pseudomonas	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella noli faenolis simbaemae impori memorine amerimoni perentinon presentences structus	Streptococcus	Salmonella
	COVERAGE	•		%	74%			101%	105%	dh
SAU102007 SeqID IDEN COVE	SeqID IDENTITY COVERAGE			11266 60% 97%				12428 100% 100%	13258 61% 97%	
SAU102032 SeqID	SeqID						12086	12198		13989
							%			75%
SAU102035		10299	10933 50%	10974 26%	11514 29%		11860 12	100%	13360 31%	13763 569
	CUVEKAGE	98%	%66	%5%			8/6	100%	98	%66
SAU102044 SeqID	SeqID	10141	10916	11011	11344		12041	414	13447	13977
	COVERAGE	00%		100%			%		102	100%
SAU102046 SeqID	SeqID	10103	<u></u>				1	12415		14001
	IDENTITY COVERAGE	32%	28%				29%	100%		29% 89%
SAU102049 SeqID	SeqID	10427	518	10962	11291		1	2416	13652	13781
	DENTITY	36%	36%	49%	40%		41%	100%	46%	36%
	COVERAGE	%[0]	%66	%16	%66		100%	100%	%86	101%
SAU102054 SeqID	SeqID		10494	8		11676	11856	4		13877
	COVERAGE	53%	50% 79%	55% 100%	51% 100%	53%	55% 100%	100%		53%
SAU102059		Г	10771	1	11622		11969	286	13226	14059
	IDENTITY COVER A GE	43%	72%	43%	40%		41%	100%	72%	40%
CA11102067 Centr	Coott	10200	10564	11156			11705	10070	12407	
100701000	DENTITY		52%	31%			- %	%00	1340/	31%
		%56		%86			%16	100%	%86	94%
SAU102068			10680					12288		
	COVERAGE		29% 101%					100%		
SAU102102								12696		
	COVERAGE							100%		
SAU102113	SeqID IDENTITY COVERAGE		10641 34% 110%)				12178 100% 101%		
								201		

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SeqID IDENTITY faccalis influenzace pylori pneumoniae aeruginosa aureus ppeumoniae typhi DENTITY 10642 1048 1048 138%	influenzae 6 85% 100% 6 98%	pylori	pneumoniae aeruginosa	veruginosa	aureus	рпештопіае	typhi
10016 10643 43% 619 101859 10859 10760 37% 39% 10154 10054 10054	885% 0010% 698%						
10016 10643 43% 619 101859 10859 10760 37% 37% 10154 10154 10154 10056 10056	85% 100% 6 98%				12180	13480	
10016 10643 43% 619 101859 60% 10760 39% 10154 37% 10154 37% 10154 32%	100% 6 98%				100% 100%	31%	
RAGE 101% RAGE 101% RAGE 10154 TTY 32% AGE 10056 TTY 32%	,000% 6 98%	11604		12027	12181	13481	13947
TTY 60% RAGE 10760 TTY 39% RAGE 10154 TTY 37% RAGE 99% TTY 32% TTY 32% RAGE 10164	%86	38%		42% 103%	100%	55%	41%
TTY 609 RAGE 10760 TTY 399 RAGE 99% TTY 32% TTY 32% AGE 100%	%86				12	3400	
TTY 399 RAGE 10154 TTY 37% AGE 10154 TTY 32% AGE 100%					100%	26%	
1TY 399 RAGE 10154 TTY 37% SAGE 99% TTY 32% TTY 32% SAGE 10054					12177	1304	
TTY 37% 3.7% 3.4GE 99% 10154 11Y 3.2% 3.4GE 100%	101%				100%	41%	
TTY 37 RAGE 10154 TTY 32 RAGE					12457		
TTY 32	_				Š		
TTY 32					100%		
RAGE					12458 100%		
					100%		
					12459		
IDENTITY COVERAGE	-				100%		
SeqID			T		12462		
IDENTITY					100%		
COVERAGE					100%		
SeqLD					12460		
COVERAGE					100%		
					12665		
IDENTITY COVERAGE			71111.1		100%		_
					12666		
IDÈNTITY COVERAGE		-		·	100%		
10447 1079	10994	11358		Π		13192	13818
COVERAGE 58% 68% COVERAGE 99% 99% 99%	%66 %66 85 %66	52% 99%		%66 63%	100%	%66 %29	28% 99%
362) 20%	11193		1	12020 38%	12527 100%	13561 46%	13731 41%
10100 10700	9370 6970		11607	74%	100%	%66	94%

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. TABLE VIIA

Data	Escherichia coli 36%	nterococcus ecalis	Haemophilus influenzae	Helicobacter Kleb pylori pneu	Klebsiella Pseudomona oneumoniae aeruginosa 35%.	Pseudomonas aeruginosa	Pseudomonas Staphylococcus Streptococcus Salmonella aeruginosa aureus preumonoccus typhi 1000 1000 1000	Streptococcus pneumoniae	Salmonella typhi
36% 75%		40% 79%			33% 74%		00% 100%	42% 79%	34% 75%
108	<u> </u>	10800 61% 98%					12531 100% 100%	13496 45% 91%	
10163 10845 28% 43 74%	<u>20</u>	1845 43% 99%							
"8	108	10847 108 72% %	953 44% 101%	1600 38% 100%	11634 11 47% 98%	1907 47% 100°	12540 100% 100%	593 70% 100%	13981 47% 100%
10274 10854 59% 74 99%	108	54 74% 100%	11154 60% 97%	11476 54% 96%		62% 62% 100	12542 100% 1009	81% 1019	13866 58% 6
							12543 13 100% 101%	180 28% 74%	
10300 10677 39% 44 79%	<u>(6</u>	77 48% 93%			11748 39% 73%	11981 37% 91%	12241 13 100% 100%		13825 41% 98%
10451 33% 97%				11515 32% 97%			12243 13 100% 101%	13531 75% 101%	
10451 38% 81%				11515 29% 75%			12244 13 100% 101%	274 85% 1019	
	1084	14 65% 97%					12245 13. 100% 100%	519 72% 979	13782 25% 87%
10182 1064 34% 96%	1064	10646 37% 87%			11682 32% 96%		12246 13. 100% 101%	275 83% 1009	13984 32% 87%
, 79%	i <u>c</u>	31 30% 80%					12247 13 100% 100%	276 74% 99	13983 26% 79%
12	12	39% 103%			11724 31% 84%		12248 100% 100%	277 82% 100	13881 34% 104%
10160 45% 100%						5103 44% 100%	12250 100% 100%		13830 43% 101%

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	Á					- 1				
LOCUSID Data	Data	escherichia coli	Escreriona Enterococcus Haemophius Heitobacter Klebstella coli faecalis influenzae pylori pneumonia	Haemophilus influenzae	Helicobacter pylori	0	Pseudomonas zeruginosa	Pseudomonas Staphylococcus Streptococcus Salmonella aeruginosa aureus pneumoniae Iyphi	Streptococcus pneumoniae	Salmonella typhi
SAU102265 SeqID	SeqID						11926	12251		1
	IDENTITY COVERAGE						37%	100%		
SAU102268								12252		
	IDENTITY COVERAGE							100%		
SATT102270	Sealth							- 1		
	IDENTITY							100%		
	COVERAGE							100%		
SAU102280	SeqID							12378		
	IDENTITY COVERAGE							100%		
SAU102281	SeqID	10316		11227	11469			12384	14.	13762
	DENTITY	45%		48%	33		45%	100%	619	44%
	COVEKAGE	- 1		%66			966	100%	100%	
SAU102283		10260 41%	10875 59%	10982 43%	11560 41%		11945 41%	12119 100%	13251 54%	14
7 00000	COVERAGE	%88	%88	%88	%26		95%	102%	%88	%88
SAU102284 SeqID	SeqID							12389		
	DENTITY COVERAGE							100%		
SAU102286 SeqID	SeqID	10385	10595						13688	
	IDENTITY COVERAGE	37% 104%	42%					100%	39%	
SAT1102287	SeaTD	10220	0594	11025		11663	11025	17308	13/177	- 1-
	IDENTITY	42%	45%	40%		%	%	%	41%	39%
	COVERAGE	81%	%56	88%		89%	84%	101%	94%	
SAU102292	SeqID	10399	10579	11018					13230	14065
	COVERAGE	41%		40%	3/%	41% 101%	42% 101%	100%	57%	41%
SAU102294	SeqID							12610		
	IDENTITY COVERAGE							100%		
SAU102297	SeqID	10405	10912	11063	11303		12117	12704	9	14066
	COVERAGE	%66			%66		%	100%	100%	48%
SAU102298 SeqID IDENTITY	SeqID IDENTITY	10404 10 36%	10914 62%	11031 33%		11686 35%	12116 28%	12705	13255 54%	
000001110	COVERAGE	72%	%66			%68	87%	100%	100%	
SAU102308 Seq.ID	SeqID	10077	10577	11248	11625	11732	12032	12706	13350	13995

Imonella	ohi 2007	39% 95%	039	31% 89%	829	ဆွဲ	95%																13960	43% 95%				13802 36% 96%
Streptococcus Sa	neumoniae typ	95% 39% 0% 100% 95%	13242 14	63%	13316 13	31%	30%												13426				324	%66 899				
Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella	aureus	87% 88% 90% 100% 45% 55% 55%	12707	100%	12657	100%	100%	12101 12658 50% 100%	100%	12659	100%	12660	100%	12655	100%	101%	12433	~	12434	100%	12435	~	9436	100%	12437	100%	12265 100% 100%	11808 12267 39% 100% 100% 100%
Pseudomonas	aeruginosa	38% 90%	90811	37%	12102	40%	%q6	12101 50%	%26						37%	%98							11805	43%			11870 32%	11808 39% 99%
Klebsiella	pneumoniae	39% 88%																										
Helicobacter Kleb.	pylori	33%																					11546	48% 98%				11386 27% 101%
Haemophilus	influenzae	37% 86%																					\sim	45% 95%				11157 111. 33% 90%
Enterococcus	faecalis	38% 46% 88% 100%	10795	75% 97%	10550	43%	%/6												10657	100%	10726	39%	10669	60% 100%				
Escherichia	coli	38%	10122	32% 90%	10057	41%	%0%	10056 50%	91%															43%				10367 36% 96%
Data		IDENTITY COVERAGE		IDENTITY COVERAGE	SeqID	,	COVEKAGE	SeqID IDENTITY	RAGE	SeqID	IDENTITY COVERAGE		IDENTITY COVERAGE	SeqID	IDENTITY	COVERAGE	SeqID	IDENTITY COVERAGE	SeqID	COVERAGE	SeqID	IDENTITY COVERAGE	SeqID	IDENTITY COVERAGE	SeqID	IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE
LOCUSID Data			SAU102318		SAU102333			SAU102334		SAU102336 SeqID		SAU102340		SAU102345 SeqID			SAU102350		SAU102352 SeqID		SAU102355 SeqID		SAU102356 SeqID		SAU102378 SeqID		SAU102380 SeqID IDEN1	SAU102388 SeqID IDENTITY COVERAGE

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LOCUSID Data	Data	Escherichia coli	Enterococcus faecalis	Haemophilus influenzae	Helicobacter pylori	Klebsiella Pseudomom oneumoniae aeruginosa	Pseudomonas aeruginosa	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Preudomonas Staphylococcus Streptococcus Salmonella coli faecalis influenzae pylori pneumoniae aeruginosa aureus pneumoniae typhi	Streptococcus pneumoniae	Salmonella typhi
SAU102389 SeqID	SeqID	10063	10547	10988			Г	12268	3	13917
	IDENTITY COVERAGE	33%	59% 97%	31%			36%	100%	35%	33%
SAU102390	1	10192				11678		12269		13753
	IDENTITY COVERAGE	41%				26% 97%		100%		42% 100%
SAU102392 SeqID		10131	10500				1951	270	13474	
	ITY	, i	42%			32%	42%	100%	42%	
	COVERAGE	73%	%08			%08	74	100%	76%	
SAU102394			10807					12271		
	IDENTITY COVERAGE		32% 102%					100%		
SAU102396 SeqID	SeqID	10243	10809					12272	ı¥	13794
	IDENTITY COVERAGE	37%	%59 86%					100%	27%	37%
SAU102401	SeqID							12209		
	DÈNTITY							100%		
	COVERAGE							100%		
SAU102417	SeqID		10934				12068	12204		
	COVERAGE		%6L 21.00					100%		
SAU102418 SeqID	SeqID					11760		12205		
	IDEN III Y COVERAGE					%68 86%		100%		
SAU102420 SeqID	SeqID							12206		
	IDENTITY COVERAGE							100%		
SAU102422 SeqID	SeqID	10308				11665	11977	12207		13776
	IDENTITY COVERAGE	30% 92%				30% 72%	27%	100%		31%
SAU102423 SeqID	SeqID			11084	17		12099	12208		
	TTY			27% 94%	25% 92%		27% 93%	27% 100% 93% 100%		
SAU102433 SeqID	7	10395 10	908	167	11616		11772	12701	13552	
	珀	101%	100	100%	73%		72%	100%	**************************************	
SAU102434	SeqID IDENTITY	10394 10	907	166 28%			11773 26%	11773 12700 134 26% 100%	46 40%	13921
	RAGE	%66	100	%66			100%	100%	101	%66
SAU102437 SeqID		10393	952	11057	11330		11774	12695	20	13920

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	Salmonella	56%	12	%66			13860	102%	14028	45%	13857	100%	13783				13917						14025 27% 97%			
	Streptococcus	64%			4	%86 %76	13435	102%	13434	100%	13237 13	100%	13265			41% 71%	13395				13475 35%	83%	13476 26% 89%			
	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella coli Itaecalis Influence influence influence inforti merimoniae periodinosi americano.	100%	<u> 2</u>	100%	12685	100%	12681	101%		97% 100%	12675	100%	1820 12674 132	100%	12669	100%	12171	100%	12172	100%	121 <i>7</i> 3 100%	100%	12174 100% 100%	12175	12405	100%
	Pseudomonas	~~	12085	∞.	!		12073 34%	101%	2072	1	12076	%66	11820	40%	12107	%67 29%	11837	Õ								
VIIA	Klebsiella Pseudomona								[73]	45% 76%			629	40% 94%												
IABLE VIIA	Helicobacter milori	51%					11332	101%	11444		11487	%66 66	11478													
	Haemophilus influenzae	57%					11049		11253	%86 6	11264 11	100%	11143		-		34%						10971 26% 105%			
	Enterococcus foecalis	%66 %L9			10947	%86 %86	10946 55%	102%	945	986	943	100	748	%86		43% 101%	10547				10868 28%	88%	10713 26% 96%			
	Escherichia coli	55%					10460	101%	10445	%2%	10456	%00I	10420	97%			10063	98%	10217	%86					10306	84%
	Data	IDENTITY COVERAGE	SeqID IDENTITY	COVERAGE	SeqID	COVERAGE	SeqID IDENTITY	COVERAGE	SeqID	COVERAGE	SeqID	COVERAGE	SeqID	COVERAGE	SeqID	COVERAGE	SeqID IDENTITY		SeqID IDENTITY	COVERAGE	SeqID IDENTITY	COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY	SeqID IDENTITY	COVERAGE
	LOCUSID Data		SAU102440 SeqID		SAU102447		SAU102448 SeqID		SAU102449 SeqID		SAUI02450		SAU102452		SAU102453		SAU102460		SAU102469		SAU102473		SAU102474	SAU102476	SAU102479 SeqID	

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LOCUSID Data	Data	Escherichia coli	Enterococcus faecalis	Haemophilus influenzae	Helicobacter pylori	Klebsiella Pseudomon pneumoniae aeruginosa	Pseudomonas aeruginosa	Escherichia Enterococcus Haemophitus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella coli faecalis influenzae pylori pneumoniae aeruginosa aureus pneumoniae typhi	Streptococcus preumoniae	Salmonella typhi
SAU102480 SeqID	SeqID	10310						12404		13770
	IDENTITY COVERAGE	28% 100%	33%				30% 100%	100%		27% 100%
SAU102481	SeqID	10289	10831					12422		13879
	IDENTITY COVERAGE	26%	29%					100%		26% 102%
SAU102485	SeqID	1	08					12421		13961
	IDENTITY COVERAGE	28%	53%					100%	%66 %95	60% 93%
SAU102486		10294	10889	11025				12420	13513	13962
			38%	%				9	42%	
20,00		95%	%16	95%				%I0I	93%	95%
SAU102487	SeqID							12419		
								100%		
SAU102498	SeqID IDENTITY	10241	10597	10974 35%	11342	37%	1842 38%	12688	13387	14092 36%
		93%		93%		94%	94%			93%
SAU102502	94 -						2060	12689		
	COVERAGE						% 82%	100%		
SAU102503							12059			
	IDENTITY COVERAGE						32%	100%	****	
SAU102526								12691		
								ò		
200	COVERAGE	600						100%		
SAU102527	SeqID	10352	09001	11104	11439		5171	12260	13204	13968
	COVERAGE	%86					93%			93%
SAU102531	SeqID		10765					12667		
	COVERAGE		34% 102%					100%		
SAU102541	SeqID	10076	10520	11000	11498		11966	12668	13405	13718
	COVERAGE	93%		91%			100%			93%
SAU102551	SeqID IDENTITY COVERAGE			11013 47% 87%	11353 38% 84%		11816 39% 84%	12672 100% 101%	13271 41% 95%	
SAU102554 SeqID	SeqID		10494					12673	13466	

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Salmonella typhi	13836 27% 98%	13859 59% 89%				13833 31% 86%	13773 32% 101%	13867 27% 92%	13971 58% 99%			13867 26% 94%	1
Streptococcus pneumoniae 44% 98%		503 73% 949			13513 27% % 88%		3 32% 77	13256 51% 97%	13200 77% 100%			256 50% 97%	13 <i>579</i> 43%
Pseudomonas Staphylococcus Streptococcus Salmonella aeruginosa aureus pneumoniae hyphi 100% 44% 98%	30%	12411 100% 101%	12537 100% 100%	12611 100% 100%	12463 13. 100% 100%	12464 100% 100%	11975 12466 13653 13773 33% 100% 32% 32% 79% 100% 100% 1011%	12467 100% 100%	12249 100% 100%	12469 100% 100%	12470 100% 100%	12471 100% 100%	12472 100%
Pseudomonas is aeruginosa o	30% 30%	12074 65% 81%		,		11979 31% 92%	11975 33% 79%	11931 12467 28% 100% 5% 93% 100%	11993 12 60% 8 99%			931 25% 93	
						11710 27% 75%		722 28% 959	1679 59% 1009			11722 27% 95%	
Helicobacter Kleb pylori pneu	11618 35% 99%	11420 51% 89%					519 30% 73%		57% 57% 100%				
Haemophilus influenzae	11232 29% 91%	11050 60% 88%				10958 32% 85%	10958 1110 30% 93%	11076 30% 93%	11100 61% 100%			11076 30% 92%	
terococcus calis 47%		948 76% 959	1		10889 27% 87%		944 26% 769		1555 78% 1009		10836 47% 96%		
Escherichia coli	10166 28% 98%	10459 10 59% 88%				10187 30% 102%	10206 36% 89%	10273 27% 95%	10356 58% 100%			10273 27% 95%	
TITY	IITY	IITY RAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	TTY	TTY	TTY	TTY	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY
LOCUSID Data	SAU102575 SeqID IDEN' COVE	SAU102578 SeqID IDENT COVE	SAU102584 SeqID IDENI COVE	SAU102585 SeqID IDENT COVE	SAU102593 SeqID IDENT COVE	SAU102598 SeqID IDENT COVE	SAU102599 SeqID IDENT COVE	SAU102601 SeqID IDENT COVE	SAU102602 SeqID IDENT	SAU102603 SeqID IDENT COVE	SAU102605 SeqID IDENT COVE	SAU102606 SeqID IDENT COVE	SAU102607 SeqID

10.11.00.11.01	typhi					13988 26% 97%	13927 32%	%68	13926 31% 100%		13881											_
Othombood	oreprococcus pneumoniae	%86									13370	59%					13696	29% 102%	13697	39%		
Redactivita Fritanciana Hammahilia Halisahadan VIIIA Decembra and Combalacana Combalacana Columba	supply occurs	100%	12473 100% 100%	12474 100%	100%	12475 100% 100%	12476 100%	100%	12477 100% 100%	12479 100% 100%	12480	100%	12481	100%	12712	100%	12650	100%	Γ	100%	12653	10.0
Decondomonage	Dneumoniae aeruginosa								12098 26% 87%						l	32%						
VIIA	preumoniae								11720 32% 92%		11724	58%			11657	44%		51 201 a 31 a				
Halicohacter VIIA	pylori																					
Hamonbilue	influenzae	•				11272 28% 95%			-													
Fatorocoone	faecalis						10600 559		10601 40% 100%		10519	62% 101%	10885 26%	108%	10522	27%						
Fecharichia	coli					10461 26% 97%	10211 33%	86%	10234 32% 98%		10288	61%										
Data		COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY	COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY	COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID	IDĖNTITY COVERAGE	SeqID IDENTITY	COVERAGE	SeqID	IDENTITY COVERAGE	SeqID	IDENTITY COVERAGE	SeqID	IDENTITY COVERAGE	SeqID IDENTITY	COVEDAGE
TOCISM Data			SAU102609	SAU102610		SAU102613	SAU102614 SeqID		SAU102615 SeqID IDENT COVE	SAU102620	SAU102621		SAU102629		SAU102631		SAU102636 SeqID		SAU102637		SAU102652 SeqID	

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					TABLE VIIA	- 1				
LOCUSID Data	Data	Escherichia coli	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella coli	Haemophilus influenzae	Helicobacter a	- 01	Pseudomonas	Pseudomonas Staphylococcus Streptococcus Salmonella aerueinosa aureus	Streptococcus nneumoniae	Salmonella tvnhi
SATTIONS SegID	Cloud	10283		Т			Т		1	13855
20701000	IDENTITY	45%	%1	47%			%	%00	49%	41%
	COVERAGE	97%					97%	100	%96	100%
SAU102663	SeqID	10304	∞	11043	9		П		13172	13780
	IDENTITY	43%	28%	44%	34%		45%	100%	. 26%	41%
	COVERAGE	%66	%66	%96	95%		9	100	97%	%66
SAU102669	Qubas	10022	1756	11257			12045	09171	371	14035
	IDENTITY	42%	76%	43%			41%	100%	24 _%	41%
	COVERAGE	%96	91%	95%			94%	100%	%56	
SAU102671	GlpaS	10409		11079	11319	683	043	191	373	14033
	DENTITY	34%		32%	44%	35%	26%	100%	%69	33%
	COVERAGE	91%		91%	%96	74	%66		%96	91%
SAU102674	SeqID	10020		11164		648	27			14016
	COVERAGE	102%		103%		101%		101%		102%
SAU102693	SeqID	10178	18		14		883	12627	301	13940
	IDENTITY	23%	74%		38%		49%	100%	61%	46%
	COVERAGE	82%	87%		%98		%98	101	%06	
SAU102694	SeqID	10177	099		11296			12628	13302	
		48%	99	20%	44%		55%	100%	60%	
	RAGE	%16	102%				(O)	1029	102%	
SAU102725		10418	10514	11137	11507					13789
		40%	72	36%	86		37%	100%	99	40%
	COVERAGE	%96	100%		103%		104%	100%	100%	- 1
SAU102764	SeqID	10179	10929	11234	11295		11884	12625	13484	13938
	COVERAGE	%66		%66			45%	100%		45% 99%
SAU102812	102		×					2127	Ĉ,	
	IDENTITY		48%	-				100%	49%	
	CUVERAGE	,	%00I					IOI	%06	
SAU102870	SeqID	10113	10880					12170	13270	14008
	COVERAGE	92%						్	93%	87%
SAU102880	SeqID	10360	533					П	13196	E
	IDENTITY COVED A GE	%09 %09	82%	61%	57%	61%	58%	100%	85%	61%
1000001	COVERAGE	8		- 1		1007	₹	10170	10100	200
SAU102881	SEQUE SENTITY COVERAGE	38%	10251 69% 98%	11099 37% 89%			38%	$12242 \\ 100\% \\ 101\%$	13199 54% 102%	38%
SATT102883 SeaTD	ContD	10396	0/0/	11168	11440		₹ Т	12702	13181	2
700701046	achre	. 02001	_	_	11447	_	_		10101	_

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	, and a second				TABLE VIIA	/IIA				
LOCUSID Data	Data	Escherichia	Enterococcus fapecalis	Haemophilus influence	Helicobacter 1	Klebsiella Pseudomono	Pseudomonas	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella coli freecilis Inthemae inflori Internation International International International Internation Internation	Streptococcus Salm	Salmonella
	IDENTITY COVERAGE	53% 86%		×	%98 %09			100%	%06 %9L	
SAU102905	SeqID IDENTITY		10732 31%	11217	11373			12273		
	_		- Se I	808	87%			100%		
SAU102909	SeqID	10042	10488	11150	457	637	940	12315	13437	13908
		95%	%56 62%	956	130%	39% 95%	86 86	100% 1019	13%	29% 95%
SAU102933	SeqID IDENTITY	10448 33%	10949	10995 35%	1579 37%	31%	985	7412 100%	13502	13817
	COVERAGE	104%		101%	108%	107%	101%	1019	101%	103%
SAU102936	SeqID	10236	10872				1804	ω.		13955
	COVERAGE	92%	100%				%96 80%	100%		33%
SAU102942	SeqID	10136	492	12		11696		552		13834
	COVERAGE	52% 100%	55%	43%		20%		100%	51%	51%
SAU102944 SeqID	SeqID					2///		12468	257	0/66
	DENTITY							100%	42%	
0.4111000000	COVERAGE	1001						100%	%66	
SAU1029/9	SeqID	33%		37%	32%		11936	12536	429	13712
		88%		87%				100%	87%	%06 06
SAU102983			10883					12676	13269	
	IDENTITY COVERAGE		28%					100% 100%	27%	
SAU102992 SeqID	SeqID			122	11297		1	12630	303	13941
	COVERAGE	%66 %70	/0% 92%	%66 %70	48%		%66 866	100%	%69 869	61% 101%
SAU103010 SeqID	SeqID							12194		
	IDENTITY COVERAGE				-			100%		<u></u>
SAU103024 SeqID	SeqID IDENTITY					11670	12042	12200		
	COVERAGE					86%	72%	101%		
SAU103025 SeqID IDENT COVE	SeqID IDENTITY COVERAGE		1,,,,			, ,,,		12202 100% 100%		
SAU103037 SeqID	SeqID		10867					12613	13267	
	COVERAGE		%66					101%	%98	

Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella Coli Geocalis Interace proprie proprie coli Pseudomonas Interace Inte
and the state of t
10936 42% 84%
10110 10783 1 43% 48% 115% 100%
101 <i>57</i> 36% 96%
10368 36% 102%
10033 10639 53% 70% 78% 80%
10372 10553 42% 74% 84% 98%
10621

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	Escherichia	Enterococcus	Haemophilus	Helicobacter Kleb	Klebsiella	Pseudomonas	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella	Streptococcus	Salmonella
0	coli	faecalis in 39% 79%	influenzae	pylori	oneumoniae aeruginosa	•	aureus 100% 101%	oneumoniae 37% 78%	typhi
' '	10259	10622	10978			12026	12720	13325	1
	73%	%26	73%			74	100%	%96 %0#	73%
$\overline{}$	10262		10984	4					14090
	51% 82%		56% 91%	57% 93%		45% .93%	100%	%89 100%	49% 82%
		10712					12734 100%		
		%66					100%		
	10109 33% 95%	10756 64% 100%	11257 34% 98%			33% 33% 95%	12739 100% 100%	13371 33% 95%	13996 32% 95%
							12751 100% 100%		
	10164 26%	584 30%	10968 25%	11566		11912	12755		13892
	%26					93%	•		%86
	10201	10478	11054			12061	12937	13425	13822
	74%	75%				81%	101%	759	74%
	10039 28% 72%	10728 31% 102%	112 <i>77</i> 26% 80%			12046 12 30% 75%	12777 100%	13423 32% 90º	13904 29% 29%
							693 100%	\	
-	66001		11170	11602	1645	11788	12780		13992
	33%		31%	31%	34% 86%	93%	100%		34%
	10098 32%		11250 34%	==		11786 12 39%	781		13 <u>991</u> 33%
	%26		%96			97%			%26
	10435 53% 99%	1 0613 73% 100%	11038 50% 99%	=		11998 12 52% 100%	12784 100% 100%	13397 64% 99	14046 52% % 99%
	10173 10 32% 92%	10856 31% 97%					2790 100% 100%	13297 29% 97	13937 34% % 94%

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LOCUSID Data	Data	Escherichia coli	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella coli faecalis influenzae pylori pneumoniae	Haemophilus influenzae	Helicobacter pylori		Pseudomonas aeruginosa	Pseudomonas Staphylococcus Streptococcus Salmonella aeruginosa aureus pneumoniae typhi	Streptococcus pneumoniae	Salmonella typhi
SAU200685 SeqID	SeqID IDENTITY							12801 100%	13185 31%	
	COVERAGE							100%	94%	
SAU200721	SeqID			11015	11541				13681	13922
	COVERAGE	40%	33%	41%	36% 94%			100%	42% 100	41% 94%
SAU200725		10118	10761	10966			11780	12933	13632	14020
	COVERAGE	, <u>8</u>	100%				- 8 `	100%	4/%	%86 8%
SAU200731	SeqID	10283	10822	ŏ				12342	13514	13855
	IDENTITY COVERAGE	55% 99%	54% 100%	44% 98%			43%	100%	51% 100%	46% 99%
SAU200740 SeqID	SeqID	10318	10554	12	11393			2798	3695	
	IDENTITY COVERAGE	48%	56%	48%	49%		50%	100%	55% 93%	48%
SAU200752	SeqID IDENTITY							12809		
								100%		
SAU200914	SeqID	10383	10714				11927	12837		13788
		%96			i	%6L 79%	%06 %/7	100%	%C7 61%	%06 807
SAU200916								12838		
	COVERAGE							100%		
SAU200928	SeqID	10439	10627	11036	11571		5179	12815	13646	14042
		86%					102%			
SAU200934			08/01					12842		13835
	IDENTITY COVERAGE	44% 72%	%66 83%				42% 82%	100%		42% 88%
SAU200949								12846		
	IDENTITY COVERAGE					,		100%		
SAU200960	SeqID IDENTITY				11500		11886	12431		
					70%		91%			
SAU200994	SeqID IDENTITY COVERAGE	10036 36% 100%	10497 62% 101%	11270 32% 100%			11865 37% 102%	12935 100% 100%	13310 35% 73%	14054 33% 99%
SAU201167 SeqID	SeqID		62.201					2887		

					HADLE VILA	WIII				
LOCUSID Data	Data	Escherichia	Enterococcus faecalis	Haemophilus influenzae	Helicobacter	Klebsiella Pseudomon pneumoniae aeruginosa	Pseudomonas zeruginosa	Escherichia Enterococcus Haemophitus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Islamonella coli faecalis influenzae pylori pneumoniae aeruginosa aureus pneumoniae typhi	Streptococcus pneumoniae	Salmonella typhi
	IDENTITY COVERAGE		37%)0% 100		
SAU201168	SeqID		10819					12889 136	13626	
	, -		102%					100%	100%	
SAU201184	SeqID	10448	10715	10995	11579		11985	807	13502	13819
	竝	40%	32% 108%				, 2	100%	111%	32% 111%
SAU201197	SeqID IDENTITY	10330 10 58%	10924 11 66%	11160	11321 53%		5215 12 58%	938 100%	49	13885 58%
	RAGE	%66	99%				%66	101	%96	
SAU201225 SeqID IDENT	ITY BAGE		812 41% 93%	33% 33%				896 100% 1004	13170 38% 87%	
SAU201236		10026	10679	11184	11613		12013	12891	505	12
	IDENTITY COVERAGE	32%	29%				% 62%	100%	30%	
SAU201301	SeqID							12899		
	COVERAGE							100%		
SAU201333	SeqID	10192				11678		12905		13753
	COVERAGE	41% 100%				%96 96%		100%		41% 100%
SAU201375	SeqID IDENTITY						11929 36% 95%	12926 100%		
SAU201380		10379	10499		11313		12024	922		13801
	IDENTITY COVERAGE	34%	25%		26%		25%	100%		25% 101%
SAU201381	SeqID	10241	10597	10974	11387	706	11833	12923	13387	13878
		%68					~~			
SAU201403	SeqID IDENTITY COVERAGE	-						12913 100% 100%		
SAU201469								12967		
SAU201486 SeqID IDEN	SeqID IDENTITY							13023 100%		
	COVERAGE							 %001	_	

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	Salmonella typhi	13841	100%	13805	30%			13996	33%	95%			14009	%96	13957	49%						13902	46%			13743	33%		
	Streptococcus pneumoniae							13625	32%	%//6	13474	73%	598	40%	13268	54%	13243	%	95%									13689 40% 92%	
	Lschertchia Enterococcus Indemoprius reitcobacter Kleostelia i Yeutomonas Siapryvococcus Streptococcus Satmonelia coli faecalis influenzae pylori preumoniae aeruginosa aureus pneumoniae typhi	12946		Įξ	100%		100%	943	100%	IOC	.942 100%	100	54	101	76	100%	973	%001	100%	12972	100%		100%	2982	100%	12981	100%	12963 100% 100%	12770
-	F seudomonas aeruginosa	11963	102%	11874	42%			2099	%	\$	11951 12	73	11875 129	%66 642.00	11905	45% 103%	11902	vo.	8	11962 12	72%	12047	47%				34% 73%		
VIIA	Klebsiella Pseudomon pneumoniae aeruginosa		******								33%	=											49%				%6L 79%		
IABLE VIIA	riencobacter pylori												11396		11357	30% 92%	11539	38%	73%			11392	42% 91%			11557	31%		
1.1	naemopnius influenzae							11257	28%	%0%			11258	94%	11213	. 4/%										11028	71%		
Ē	Enterococcus faecalis									00.00	39%				951	61% 94%						10842	53%			10900		10623 45% 89%	
7 2	coli	10145	101%	10370	73%	10229	29%	10109	33%	92%	50%	71%	10112	%,	10224	%86 %0¢						10038	49% 91%			10291	71%		
7	Data	SeqID	COVERAGE	SeqID	RAGE		COVERAGE	SeqID	IDENTITY Cover ACE	COVERAGE	Sequi		SeqID IDENTITY	COVERAGE	SeqID	COVERAGE	SeqID	IDENTITY	COVERAGE	SeqID IDENTITY	COVERAGE		IDENTITY COVERAGE		IDENTITY COVERAGE	SeqID		SeqID IDENTITY COVERAGE	SeqID
T OCTION	TOCOSID	SAU201506 SeqID		SAU201508		SAU201513 SeqID		SAU201539 SeqID		1 4 7 7 0 0 1 7 4 7	SAU201341		SAU201558		SAU201571		SAU201611			SAU201615		SAU201621		SAU201654		SAU201666 SeqID		SAU201752	SAU201765 SeqID

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almonella	ingvi				14088 39% 108%						14085 52% 94%			
Streptococcus	еитопае				411 63% 101%					13374 48% 98%	417 58% 94%			
Escherichia Enterococcus Haemophilus Helicobacter Kiebsiella Pseudomonas Staphylococcus Streptococcus Salmonella	aureus pr 100% 100%	12996 100% 100%	12996 100% 100%	12769 100% 100%	13002 100% 100%	13008 100% 100%	13020 100% 100%	13015 100% 101%	13018 100% 100%	13009 13 100% 101%	714 100% 101%	12895 100% 101%	12895 100% 101%	12731 100%
Pseudomonas	pneumoniae aeruginosa				11787 45% 88%						11946 12 46% 93%			
Klebsiella	рпеитопіае													
Helicobacter Kleb	pytori				11310 41% 104%					11359 44% 96%	⊨			
Haemophilus	ınfiuenzae				11134 41% 100%						10983 52% 91%			
Enterococcus	jaecalis				10783 46% 100%						10874 50% 94%			
Escherichia	1100				10258 38% 108%						10261 51% 94%			10062 28%
Data	IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY COVERAGE	SeqID IDENTITY
LOCUSID Data		SAU201773	SAU201775 SeqID IDENT COVE	SAU201810 SeqID IDENT COVE	SAU201827	SAU201929	SAU201952	SAU201971 SeqID IDENT COVE	SAU202006 SeqID IDENTITY COVERAG	SAU202039 SeqID IDENTITY COVERAG	SAU202126 SeqID IDENT COVE	SAU202174 SeqID IDENT COVEI	SAU202176 SeqID IDENT COVE	SAU202186 SeqID IDENTITY

LOCUSID Data	Data	Escherichia	Enterococcus	Haemophilus	Helicobacter Kleb.	Klebsiella	Pseudomonas	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella	Streptococcus	Salmonella
		coli	faecalis	influenzae	pylori	pneumoniae aeruginosa		aureus	pneumoniae	typhi
	COVERAGE	73%						101%		
SAU202267 SeqID IDEN' COVE	SeqID IDENTITY COVERAGE							12727 100% 100%		
SAU202708 SeqID	SeqID	10428	10913					12855		13735
	IDENTITY	25%	28%	_				100%		25%
SAU202736 SeqID	SeqID	10148	10902	11181	11494	11677	11857	12927	13248	13844
	IDENTITY COVERAGE	39%		%86	%16	%08 %08	38%	100%	38%	39%
SAU202756 SeqID	SeqID	10436	10614	11071				13027	13246	14045
	IDENTITY	44%	63%	47%	•		44%	100%	53%	40%
SAU202781								12718		
	IDÉNTITY. COVERAGE						-	100%		
SAU202872			10656					12866	13670	
	DENTITY		45%					100%	28%	
	COVERAGE		101%					100%	%86	
SAU202882								12848		
	COVERAGE							100%		
SAU202930	,							12871		
	IDENTITY COVERAGE							100%		
SAU202945 SeqID	SeqID							12868		
	IDENTITY							8		
	COVERAGE							100%		
SAU202968 SeqID	SeqID							12886		
	IDENTITY COVERAGE							100%		
SAU203001	SeqID							12894		
	IDÊNTITY				_			100%	-	
	COVERAGE							100%		
SAU203007	SeqID IDENTITY							12893 100%		
	COVERAGE							100%		

					TOTAL VILLA	אווע				
LOCUSID Data	Data	Escherichia coli	Enterococcus faecalis	Haemophilus influenzae	Helicobacter pylori	Klebsiella Pseudomon pneumoniae aeruginosa	Pseudomonas aeruginosa	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella coli (aecalis influenzae pylori pneumoniae aeruginosa aureus pneumoniae lyphi	Streptococcus pneumoniae	Salmonella typhi
	SeqID IDENTITY COVERAGE							12945 100% 101%		
SAU203293	SeqID IDENTITY COVERAGE					1		12979 100% 101%		
SAU203296	SeqID IDENTITY COVERAGE				11330 29% 88%			12263 100% 101%		
SAU203524	SeqID IDENTITY COVERAGE							00%		
SAU300110	SeqID IDENTITY COVERAGE	10054 33% 82%	10544 38% 109%			11662 33% 73%		13031 100% 102%	13441 30% 109%	
SAU300131	SeqID IDENTITY COVERAGE	10344 45% 100%	10	11112 44% 100%	11434 52% 99%		5164 47% 99%	13034 100% 101%	13213 60% 99%	14103 44% 100%
h	SeqID IDENTITY COVERAGE							00%		
	SeqID IDENTITY COVERAGE		10562 43% 103%		11519 39% 91%		11844 32% 72%	12367 100% 101%	13522 41% 104%	
	SeqID IDENTITY COVERAGE				11522 32% 108%			12717 100% 100%		
SAU300617	SeqID IDENTITY COVERAGE		10851 50% 97%					12513 100% 100%	13289 49% 97%	
SAU300713	SeqID IDENTITY COVERAGE		10767 26% 83%				11823 30% 93%	13058 100% 100%		
SAU300719	SeqID IDENTITY COVERAGE	10468 46% 100%	10	11246 34% 101%	11380 30% 94%	11644 30% 101%	11887 40% 100%	12987 100% 101%	13456 33% 96%	13726 34% 100%
SAU300732 SeqID IDEN COVE	SeqID IDENTITY COVERAGE	10282 10 26% 71%	10682 51% 88%					00% 100%	13394 49% 86%	
czennenwe	Ozdar.	_	1002	_		_	_	15008	170/1	_

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ä	LOCUSID Data	Escherichia	Enterococcus faecalis	Haemophilus influenzae	Helicobacter pylori	Klebsiella Pseudomon pneumoniae aerueinosa	seudomonas	Escherichia Enterococcus Haemophitus Helicobacter Klebsiella Pseudomonas Siaphylococcus Streptococcus Salmonella coli faecalis influenzae invlori pneumoniae aerueinosa aureus pneumoniae tynhi	Streptococcus presumoniae	Salmonella tvohi
	IDENTITY COVERAGE		52%					100%	41% 97%	
SAU300975	SeqID IDENTITY COVERAGE		10604 30% 72%				-	12203 100% 102%		
8660	SAU300998 SeqID IDENTITY COVERAGE		10820 40% 99%					13077 100% 102%	13489 40% 99%	
SAU301004			10744 40% 101%					13079 100% 100%		
SAU301030	SeqID IDENTITY COVERAGE							13080 100% 100%	:	
0801	SAU301080 SeqID IDENTITY COVERAGE							13083 100% 100%		
SAU301118	SeqID IDENTITY COVERAGE	10242 47% 98%	10808 58% 98%	11092 48% 91%		11653 53% 78%		12904 100% 100%		13795 48% 96%
SAU301133	SeqID IDENTITY COVERAGE		%96 %					13087 100% 100%	443 30% 93%	
SAU301223	SeqID IDENTITY COVERAGE	10297 10 31% 104%)640 , 50% 99%	10964 31% 102%	11323 32% 90%		11783 13 34% 102%	3090 100% 100%	664 48% 98%	13737 32% 104%
1230	SAU301230 SeqID IDENTITY COVERAGE	10252 52% 95%	877 52% 92%	11010 63% 74%		11669 52% 95%	11956 13 59% 77%	13092 13 100% 100%	506 59% 92%	13704 52% 95%
1268	SAU301268 SeqID IDENTITY COVERAGE							.102 100% 100%		
1275	SAU301275 SeqID IDENTITY COVERAGE	10048 54% 99%	10926 47% 84%	11014 55% 97%	11511 50% 97%		11934 II. 53% 97%	3103 100% 1019	366 46% 84%	13897 54% 99%
SAU301357	SeqID IDENTITY COVERAGE		10696 74% 98%	11063 32% 80%		11766 33% 93%		100% 100% 101°	13354 76% 100%	
1433	SAU301433 SeqID IDENTITY COVERAGE							12845 133 100% 100%	13393 26% % 91%	

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					IABLE VIIA	VIIA				
LOCUSID Data	Data	Escherichia coli	Enterococcus faecalis	Haemophilus influenzae	Helicobacter pylori	Klebsiella Pseudomom pneumoniae aeruginosa	Pseudomonas reruginosa	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella coli faecalis influenzae pylori pneumoniae aeruginosa aureus pneumoniae typhi	Streptococcus pneumoniae	Salmonella typhi
SAU301465 SeqID	SeqID	10210	999	11214			11921	13013	13418	13925
	IDENTITY COVERAGE	29%	54% 104%	32% 104%	37% 100%		28% 101%	100%	52% 103%	29% 102%
SAU301472	SeqID IDENTITY COVERAGE	10157 36% 85%						12925 100% 100%		
SAU301592	SAU301592 SeqID IDENTITY COVERAGE							13137 100% 100%		
SAU301620	SeqID IDENTITY COVERAGE							13140 100% 100%		
SAU301758 SeqID IDEN COVE	SeqID IDENTITY COVERAGE							13156 100% 100%		
SAU301773	SeqID IDENTITY COVERAGE							12729 100% 100%		
SAU301829	SeqID IDENTITY COVERAGE	10107 45% 98%			11309 40% 97%		11857 42% 96%		13248 38% 106%	13935 41% 99%
SAU301869	SeqID IDENTITY COVERAGE		10732 30% 80%		11373 36% 95%	-		12903 100% 100%		
SAU301898	SAU301898 SeqID IDENTITY COVERAGE		10932 27% 71%					130 <i>57</i> 100% 100%		
SAU302060 SeqID IDENT COVE	SeqID IDENTITY COVERAGE							13042 100% 100%		
SAU302513	SeqID IDENTITY COVERAGE							12851 100% 100%		
SAU302626	SeqID IDENTITY COVERAGE							13105 100% 100%		
SAU302685 SeqID IDEN' COVE	SeqID IDENTITY COVERAGE							13113 100% 10725		
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LOCUSID Data	Data	Escherichia	Enterococcus	Haemophilus	Helicobacter	Klebsiella	Pseudomonas	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Samonella	Streptococcus	Salmonella
-		coli	faecalis	influenzae	pylori	pneumoniae	pneumoniae aeruginosa aureus	aureus	pneumoniae typhi	yphi
	IDENTITY						1	100%		
	COVERAGE							%001		
SAU302699 SeqID	SeqID							13115		
	IDENTITY							100%		
	COVERAGE							100%		-
SAU302805 SeqID	SeqID				11345			13133		}
	IDENTITY				33%			100%		
	COVERAGE				75%			101%		
SAU302901 SeqID	SeqID							12872		
	IDENTITY							100%		
	COVERAGE							100%		
SAU302931 SeqID	SeqID							13155		
	IDENTITY							100%		
	COVERAGE							100%		
SAU302950 SeqID	SeqID							12664		
	IDENTITY							100%		
	COVERAGE							101%		
SAU302956 SeqID	SeqID	10023		11256		11742	12044	12930	13372	14018
	IDENTITY	32%		28%		31%	76%	100%	31%	32%
	COVERAGE	%88		%88		%88	%98	101%	%88	%88

Data Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella controlis influence information production information production in the control of	Escherichia Enterococc	Enterococc faecalis	Sm.	Haemophilus .	Helicobacter I	Klebsiella Pseudomon	Pseudomonas	Staphylococcus	Streptococcus	Salmonella
3	3		11256			11742	Т	ano ano		14018
100			%99			95%	92%		41%	97
KAGE	000%			98%		100%	%66		%/6	100%
<u> </u>	10052	-			11503			12626		13932
IDENTITY 100% COVERAGE 100%	100%				41%		48%	38%		40%
10064	10781	10781	10993		11499		11959	12884	13614	13915
TTY 100% 50%	% 20%	%(7	%	38%		71%	45%	%	94%
COVERAGE 100% 96%				100%	%16		%16	%16	%16	%66
10065 10653 1099	10653 1099	109	10992		11311		11958	12883	13177	13916
100% 53%	53%	53%		81%	46%		71%	57%	20%	%86
COVERAGE 100% 95%	100%			101%	98%		97%	92%	92%	100%
Seq ID 10120 IDENTITY 100%	10120 100%					11768 72%				
COVERAGE 100%						85%				
10214 10608 11112	10608	1112	11129				11852			13931
100% 36%	36%	36%		74%		94%	36%		36%	%96
%96 %001	%96 %001	%96		100%		100%	97%		%16	73%
10228	1120	1120	1120	4		11631		13132		13963
100%			•	45%		%98	51%	35%		%/28
COVERAGE 100%				100%		%18	100%	100%		100%
10.	10247						11812			13948
<u>8</u>	%00I						47%			%66
COVERAGE 100%	001						93%			%96
10258 10628 1113	110628 1113	1113	11134		11489		5192	12526	13636	14088
IDENTITY 100% 46% 7	46%		1	71%	48%		71%	25%	47%	%26
COVERAGE 100% 101%				100%	100%		100%	100%	82%	100%
10266 105	10510		1126	6	11524		61811	12915	13279	14049
100% 51%	51%	51%		76%	30%		28%	45%	49%	86%
COVERAGE 100% 93%				%08	94%		%16	%96	101%	%66

- 1					TABLE VIIA	¥				
LOCUSID	Data	Escherichia	Enterococcus	Haemophilus	Helicobacter .	Klebsiella	Pseudomonas	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Sdimonella	Streptococcus	Salmonella
		coli	faecalis	influenzae	pylori	oneumoniae aeruginosa		anreus	pneumoniae	typhi
ECO103101	Seq ID	10315	10763				12052		13662	13764
	IDENTITY COVERAGE	100%	37%	73%	26%	96% 100%	64%		33%	94%
ECO104120	Seq ID	10462	10609	11034		11726	11853			13887
	IDENTITY	100%	73%	34%		87%	78%			37%
	COVERAGE	100%	%61	%68		100%	%68			%76
ECO104268	Seq ID		10607							13707
	IDENTITY	100%	43%					43%	38%	95%
	COVERAGE	00%	%26					%66	92%	100%
KPN100432	Seq ID		10736					12789		14088
	IDENTITY COVER A GE	90%	37%	62%	37%	100%	62%	41%	47%	92%
KPN100854	Sen ID	10086	10652	11107	11565	11630	11862		13380	14060
	IDENTITY	35%	29%	26%	~	7001	42%		%	35%
	COVERAGE	74%				100%			71%	74%
KPN101022	Seq ID	10475	10607			11642				13707
	IDENTITY	%06	75%			100%		27%	76%	%16
	COVERAGE	100%	77%			101%		101%	79%	101%
KPN101026	Seq ID IDENTITY	10228 86%		11204		11631	12038	13132 37%		13963
	COVERAGE	%66		%1.6		100%				%66
KPN101729	Seq ID							13032		
	COVERAGE			50% 96%	%96 6%	100%	%96 8%	63%		
KPN101750	Seq ID	10052			11503	11652	12078	12626		13918
	IDENTITY COVERAGE	94%			38% 103%	%00 %00	47%	37%		34%
KPN102057	Seq ID	10406	10892	11035		11661	11854	13153		13883
	IDENTITY COVERAGE	%6Z 6%	30%	30%		100%	27%	28% 85%		29%
KPN102638	Seq ID		10510			11667		12915		14049
	COVERAGE	77%	51% 79%		29% 83%	100% 100%		44%	50% 79%	77%
KPN103882	Seq ID IDENTITY	10315	10763	11215	11454	11716	12052		13662	13764
	COVERAGE	100%				100%			74%	101%

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COCUSID	Data	Escherichia	Enterococcus	Haemophilus	Helicobacter	Klebsiella	Pseudomonas	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus	Streptococcus	Salmonella
		coli	faecalis	influenzae	pylori	pneumoniae	pneumoniae aeruginosa	aureus	niae	typhi
XPN104183	Seq ID	10065	10653	10992	11490	11650	11958	12883		13916
	IDENTITY	97%	26%	%08	46%	<u>5</u>	%0%	%09	25%	%86
	COVERAGE	85%	74%		%98			74%	74%	82%
KPN104281	QI bəs	10023		11256		11742	12044			14018
	DENTITY	95%		%89		100%	%99		41%	%56
	COVERAGE	94%		%76		101%	94%		%16	101%
KPN104538	QI baS	10462	10609	11034		11726	11853			13887
	IDENTITY	87%	27%	32%		100%	29%			38%
	COVERAGE	100%	87%	%68		100%	%68			94%
KPN104716	CII bəS	10214	10608	11129		11757	11852		13627	13931
	IDENTITY	94%	36%	75%		100%	36%		35%	94%
	COVERAGE	100%	%96	100%		100%	%/6		97%	73%
KPN105779	Seq ID					11770	12103			
	DENTITY					100%	78%			
	COVERAGE					101%	%66			
XPN106659	Seq ID			10993		11649		12884		13915
	IDENTITY	%06	28%	72%		100%	74%	21%	28%	%16
	COVERAGE	80%	20%			101%	74%	72%		%18
KPN106840	QI bəs	10259	10857	10978				12182		14087
	IDENTITY	91%	44	74%		100%	25%	38%	42%	%16
	COVERAGE	100%	101%	%86		100%	%66	94%	92%	100%
XPN107776	Seq ID	10222		11132		11771	11810			13936
	IDENTITY	78%		37%		100%	35%			%08
	COVERAGE	98%		%68		102%	87%			%86
SAU100968	Sed ID	10064		10993	11499		11959	12643	13614	13915
	IDENTITY	45%	62%	44%	36%		46%	100%	62%	46%
	COVERAGE	97%	%16	100%	%66		97%	100%	%86	%16
SAU201145	QI bəs		10781	10993	11499		65611	12884	13614	13915
	IDENTITY	45%	62%	44%	36%		46%	100%	62%	46%
	COVERAGE	97%	%26	100%	%66		%16	100%	%86	%16
SPN101971	CII bəs	10064		10993	11499			12884		13915
	IDENTITY	46%	77%	42%	36		48%	62%	100%	46%
	COVERAGE	100%	%66	102%			100%		100%	100%
SPN201024	Seq ID IDENTITY	10064	10781	10993	11499 36%		11959 49%	12884	13614	13915
	COVERAGE	%66					%66			%66

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TOCUSID	Data	Escherichia	Enterococcus	Haemophilus	Helicobacter	Klebsiella	Pseudomonas	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella Pseudomonas Staphylococcus Streptococcus Salmonella	Streptococcus	Salmonella
		coli	faecalis	influenzae	pylori	pneumoniae aeruginosa		aureus	pneumoniae	typhi
STY000277	Seq ID	10475	10901					12370	13166	13707
	IDENTITY	%56	44%					42%	38%	100%
	COVERAGE	100%	91%					%66	%96	100%
STY000625	Seq ID	10421								13784
	IDENTITY	63%								100%
	COVERAGE	100%								101%
STY000773	Seq ID	10315	10763	11215	11454	11716	12052		13662	13764
	IDENTITY	94%	36%	71%	79%	93%	62%		31%	100%
	COVERAGE	100%	74%	100%	77%	100%	100%		74%	100%
STY001430	Seq ID	10064	10781	10993	11499		11959	12884	13614	13915
	IDENTITY	94%	46%	70%	37%		20%	46%	47%	100%
	COVERAGE	100%	%96	101%	%86		%86	%16	%86	100%
STY001433	Seq ID	10065	10653	10992	11311		11958	12883	13177	13916
	IDENTITY	%86	53%	82%	46%		72%	28%	20%	100%
	COVERAGE	%66	94%	100%	%16	•	%66	94%	94%	100%
STY001867	Seq ID	10247					11812			13948
	IDENTITY	%66					47%	•		100%
	COVERAGE	%86					%96			100%
STY002995	Seq ID	10023		11256		11742	12044		13595	14018
	IDENTITY	%26	-	%19		%56	65%		40%	100%
	COVERAGE	94%		%26		101%	94%	Ì	%16	101%
STY003357	Seq ID	10228		11204		11631	12038	13132		13963
	IDENTITY	%28		42%		85%	46%	36%		100%
	COVERAGE	100%		100%		81%	101%	100%		100%

Salmonella typhi		13899 95% 28%	14048 91% 67%		13778 77%	29%	14047	%08 %		13880	%66		14034	14034 1019	4	14034 1019 13730	14034 1019 13730	14034 1019 13730 95%	14034 1019 13730 95%
Streptococcus pneumoniae						,,,,,,	13316 91%	34%		13281	73%		13511	13511 96%	1351	96%	13511	96%	96%
Pseudomonas Staphylococcus Streptococcus aeruginosa aureus pneumoniae			12844 94% 36%		12781 96%	28%	12375 96%	38%		12351	72%		12159	12159 100%	1215	12159	12159	12159	12159
Klebsiella Pseudomonas pneumoniae aeruginosa	5053 100% 100%	5054 100% 100%	5055 100% 100%	5056 100% 100%	5057 100	100%	2028 100%	100%	5059 100% 100%	2060	100%	ı	Tonc	1000	<u> </u>	1000	1009	5062 1009	5062 1009
Klebsiella pneumoniae					11701 83%	28%						11740		11747 101% 74%	101% 74%	101%	101%	101%	101%
Helicobacter pylori			11388 91% 32%		11386 77%		11466 99%	79%				11307	1771	88%	1	88%	88%	88%	88%
Haemophilus influenzae	:	10959 94% 28%			11250 73%	32%				11275	72%	11088	20011	<u> </u>	100%	2	2	2 1 %	20 1 25
Escherichia Enterococcus Haemophilus Helicobacter Klebsiella coli faecalis influenzae pylori pmeumonia						0.00	0001 97%	35%		10785	73%								
Escherichia coli		10386 96% 28%	10265 93% 67%				10264 100%	81%		10278	%66 //3%	10408		%16		97%	97% 10324	97%	97% 10324 94%
	SeqID COVERAGE IDENTITY	SeqID COVERAGE IDENTITY	SeqID COVERAGE IDENTITY	SeqID COVERAGE IDENTITY	SeqID COVERAGE		Ή	DENTITY	SeqID COVERAGE IDENTITY		COVERAGE	SealD		COVERAGE	COVERAGE IDENTITY	COVERAGE IDENTITY SeaID	COVERAGE IDENTITY SeqID	COVERAGE IDENTITY SeqID	COVERAGE IDENTITY SeqID COVERAGE
LOCUSID Data	PA0028	PA0120	PA0129	PA0141	PA0221	2,000	FA0265		PA0321	PA0337		PA0353				PA0378	PA0378	PA0378	PA0378

LOCUSID Data	Escherichia	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella	Haemophilus	Helicobacter		Pseudomonas	Pseudomonas Staphylococcus Streptococcus	- 1	Salmonella
coli	•	faecalis	influenzae	pylori	ē	aeruginosa	aureus		typhi
10078	78	10858					12993	13560	13723
	99% 26%	100% 31%				100%	%EE 33%	100%	99% 26%
						5064			
						100%			
1						5065			
						100%			
- 13						100%			
<u> </u>	10296 100% 46%	10871 93% 79%	11003 102% 45%		11660 78% 47%	5066 100% 100%	12971 100% 27%	13461 91% 20%	13738 100% A7%
12	10123			11424		2067	708		14038
	%66			%26		%0	75%		%66
	75%			32%		100%			%91
ı						2068			
					-	100%			
12	10471					5069			
	88%				-	100%			
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\circ	10150		11237 83%	11581		5073	12153	13459	13846
	38%		38%			100%	ŧ		39%
						5074 100%			
						100%			

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LOCUSID Data	Data	Escherichia coli	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella coli faecalis influenzae pylori pneumonia	Haemophilus . influenzae	Helicobacter pylori	g,	Pseudomonas , aeruginosa	Pseudomonas Staphylococcus Streptococcus aeruginosa aureus pneumoniae		Salmonella typhi
PA0788	SeqID			1			5075			
	DENTITY						100%			
PA0882	SeqID COVERAGE	10233					5076 100%			14013 101%
	IDENTITY						%			28%
PA0934	SeqID COVERAGE	10276 1019	10876 93%	11006 1019		80%	100%	12646 92%	13483 94%	
00000	DENIIIY	4./%	40%	40%		37%	%00I	39%	38%	
PA0938	SeqID COVERAGE IDENTITY						5078 100% 100%			
PA1019	SeqID	104	10592	11180			5079			
	DENTITY	26%	25%	28%	_		100% 100%			
PA1072	SeqID COVERAGE	10377					5080		13410	13813
	DENTITY						100%		36%	
PA1115	SeqID						5081			
	DENTITY		,				100%			
PA1270	SeqID	10328				11751	5082			13946
	IDENTITY					%				76%
PA1301	SeqID	Ξ.					2083			
	COVERAGE	96%					100%			
PA1360	SeqID	10104					5084			14000
	COVERAGE	92% 63%					100%		97%	92%
PA1365	SeqID						5085			
_	COVERAGE						100%			
PA1398	SeqID COVERAGE						5086 100%			
	IDENIIIY						100%			

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LOCUSID Data	Data	Escherichia	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella	Haemophilus	Helicobacter		Pseudomonas	Pseudomonas Staphylococcus Streptococcus	Streptococcus	Salmonella
		coli	faecalis	influenzae	pylori	pneumoniae aeruginosa	- 1	aureus	pneumoniae	typhi
PA1462	SeqID		10915		11559		2087			
	COVERAGE	_	%86		101%		100%			
	DENTILI		02.67		30%		100%			
PA1493	SeqID	<u>5</u>					2088			13786
	COVERAGE	%7.6				%/.6	3			92%
	IDENTILY	30%				49%	100%			20%
PA1547	SeqID				11377		5089			
	COVERAGE	_			%88		100%			
	IDENTITY				28%		100%			
PA1636	SeqID	<u>≅</u>					2060	12990		13890
	COVERAGE	101%					%001	%96		81%
	IDENIIIY	3/%				- 1	100%	26%		32%
PA1684	SeqID					11693	5091			
	COVERAGE					%66	100%			
	IDENTITY					26%	100%			
PA1868	SeqID	103					5092			
	COVERAGE	82%					100%			
	IDENTITY	35%					100%			
PA1876	SeqID						5093			14036
	COVERAGE					%92	100%			93%
	DENTITY					40%	100%			39%
PA1918	SeqID	101		11033			5094			13745
	COVERAGE	79%		82%			100%			462
	IDENTITY	31%		28%			100%			28%
PA1986	SeqID						5005			
	COVERAGE						100%			
	IDENTITY						100%			
PA2009	SeqID						9609			
	COVERAGE						100%			
	IDENTITY						100%			
PA2083	SeqID	<u>2</u>				11692	2097			
	COVERAGE	87%				85%	2			
١	IDENTITY	31%				35%	100%			
PA2101	SeqID COVERAGE	10198 92%					5098		13282	13861
	IDENTITY						100%		25%	
										7

Salmonolla	typhi	13996 96%	37%	13893	33%	13985	%65 %86	3852	99%	13830	100%	73%										-			13930	- %86 - %86	%09	13980	29%
Г			29%		80% 27%	I						-					•		_							4			
Prendomonas Stanhalococus Strentococus	aureus	12943 94%	34%							12917	%16	44%																	
Pseudomonds	aeruginosa	5099 100%	100%	5100		5101	100%	5102	100%	5103	100%	100%	5104	100%	2001	100%	100%	5106	100%	100%	5107	100%	5108	100%	5109	100%	100%	5110	100%
	ø				80% 25%														****										
Recharichia Enterococcus Haemanhilus Helicoharter Klebsiella	pylori																												
Hoemonhilus	influenzae	112 <i>57</i> 95%	27%																									11172	28%
Fatorocous	faecalis			10865																									
Recharichia	coli	10109 96%	37%	10472	27%	10181	%09 %86	10169	99%	10160	100%	74%						10132	%98 %98	35%								10189	32%
		SeqID COVERAGE	IDENTITY	SeqID	DENTITY	SeqID	COVERAGE		COVERAGE	SeqID	COVERAGE	IDENTITY	SeqID	COVERAGE IDENTITY	Coott	COVERAGE	IDENTITY	SeqID	COVERAGE	IDENTILY	SeqID	DENTITY	SeqID	COVERAGE	SedID	COVERAGE	TTY	SeqID	IDENTITY
TOCTISTO Data		PA2108		PA2128		PA2147		PA2196		PA2197			PA2222		DA7212			PA2398			PA2424		PA2461		PA2470			PA2488	

	streptococcus salmonetta pneumoniae typhi	13719		140	92% 94% 42% 58%						13959	95%	80%			14029	101%	4270					13302	97%			13848	13848
	r seuaomonas Staphytococcus Istreptococcus aeruginosa aureus pneumoniae				99%									_									3628	97%			10000	75%
	r seuaomonas aeruginosa	5111	100%	5112	100%	5113	100	100%	5114	100%	5115	100%	100%	3116	100%	5117	100%		5118	100%	5119	100%	5120	100%	5121	100%	1007	100%
ABLE VIIA	heosieita rseudomon pneumoniae aeruginosa					11714	%08	45%								11730	90%	9/ C+										
IAB	neucooacier pylori	11516		11504	97%																		11296	89% 47%			20011	%98 86%
1.1	паеторишь influenzae	11145 98%	31%	29601	94%																		11222	84%			11922	100%
5	eoli faecalis influenzae pylori pneumonia			66801	99%						10566	%68				10703	74%						10660	%1%			10,605	79%
1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1	coli	10331 99%	42%		94%	10116	%26	41%	10441	7.7%		95%	9070			10444	101%	- 1	10384 99%	33%			10177	91%			10151	%0
	Data	用	IDENTITY		COVERAGE	SeqID	COVERAGE	IDENTITY	SeqID	DENTITY		COVERAGE	DENIII I	Sequin Co.	DENTITY	SeqID	COVERAGE		SeqID COVERAGE	IDENTITY	SeqID	COVERAGE	П	COVERAGE	SeqID	COVERAGE	Т	SAGE
TOTION		PA2494		PA2584		PA2594			PA2634		PA2641		10000	FA20/1		PA2680			PA2684		PA2726		PA2742		PA3006		DA 2011	11000

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LOCUSID Data	Data	Escherichia	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella	Haemophilus	Helicobacter	Klebsiella	Pseudomonas	Pseudomonas Staphylococcus Streptococcus	Streptococcus	Salmonella
		coli	faecalis	influenzae	pylori	pneumoniae	aeruginosa	aureus	pneumoniae	typhi
PA3013	SeqID	107	10494	11095	11525			12461		13750
	COVERAGE	98%	80%	102%	102%		100%	102%		98%
DA 2041	Charle	10207					1013	7		12777
r A3041	Seque						1000			13///
	DENTITY						100%			32%
PA3048	SealD	10117		99601			5125			14005
	COVERAGE			75%			100%			%66
	IDENTITY	47%		45%			100%			47%
PA3068	SeqID						5126			
	COVERAGE						100%			-
	DENTITY						100%			
PA3121	SeqID	<u>8</u>		11164	11363		5127	15		14017
	COVERAGE	666		%66	81%		100%	%66		%66
	IDENTITY	63%		29%	79%		100%	26%		62%
PA3153	SeqID						5128			
	COVERAGE						100%	_		
	IDENTITY						100%			
PA3154	CIpeS						5129			
	COVERAGE	_					100%			
	DENIIIY						100%			
PA3160	SeqID						5130			
	COVERAGE						100%			-117
	IDENTITY						100%			
PA3279	SeqID						2131			
	COVERAGE						100%			
	DENTITY						100%			
PA3280	SeqID						5132			
	COVERAGE	_					100%			
	DENTILY						100%			
PA3374	SeqID	5					5133			
	COVERAGE	%66 					100%			
	IDENTITY	25%					100%			
PA3479	SeqID						5134			
	DENTITY	_					100%			

TOCHEM	Data	Recharichia	Feelwairin Fatarocon Hamonilus Holiophatan Klahaialln	Hamonbilue	Halinohantar	- [Demidomonde	Desudomonae Gembuloscome Chantococus	Chrantococcus	Calmonalla
Trocosm Trocosm	Data	coli	faecalis	influenzae	pylori	g	aeruginosa	aureus	pneumoniae	typhi
PA3984	SeqID	<u> </u>		11002		1	5147			14061
	COVERAGE			98% 37%		91% 39%	100%			99%
PA4024	SeqID	102	10700			11736	5148			13951
	DENTITY	%05 20%	%05 20%			72%	100%			%05 20%
PA4027	SeqID						5149			
	DENTITY						100%			
PA4037	SeqID	101					5150	12958	13296	14002
	COVERAGE	72%	83% 30%	72% 33%	72% 34%	72% 33%	100%	70%	71%	72%
PA4067	SeqID	101					5151			13845
	COVERAGE	98%					100%			99%
PA4070	SedID	10159					6315			
	COVERAGE						100%			
	IDENTITY	28%					100%			
PA4081	SeqID						5153			
	COVERAGE	_					100%			
	IDENIII I						100%			
PA4105	SeqID						5154			
	DENTITY						100%			
PA4124	SeqID						5155			14023
	COVERAGE						100%			93%
	IDENTITY						100%			64%
PA4125	SeqID	_					5156			14024
	COVERAGE						100%			94%
PA4158	SeqID	10080	10610	11009	11379	11769	5157	12297		13725
	COVERAGE	%86	95%	88 %	83%	74%	100	%96		%26
	IDENTITY	%19	38%			61%	100%	20%		%19
PA4237	SeqID COVERAGE	10333 91%	10542	11123 98%	11582		%001 100%	12232	13224	14093
,	IDENTITY				43%		100%	45%	429	7.7%

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LOCUSID Data	Data	Escherichia	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella	Haemophilus	Helicobacter		<u> </u>	Pseudomonas Staphylococcus Streptococcus	Streptococcus	Salmonella
		coli	faecalis	influenzae		pneumoniae	aeruginosa	aureus	рпеитопіае	typhi
PA4242	SeqID COVERAGE	10338 100%	10538	11117	11428 100%		5159 100%			
	IDENTITY						100%		:	
PA4244	SeqID	10340	10534	11116			5160	12225	13217	14099
	COVERAGE	100%	100%	100%			100%	100%	100%	100%
	IDENTITY	%59	46%	%69			100%	45%	43%	92%
PA4245	SeqID	10341	10532	11115			5161	12223	13216	13812
	COVERAGE	95%	%86	%56			100%	%86	%86	78%
	IDENTITY	26%	45%	28%			100%	45%	40%	33%
PA4246	SeqID	10342	10531	11114	11432			12222	13215	14101
	COVERAGE	100%	92%	%66	%88		100%	%66	92%	100%
	IDENTITY	77%		74%	46%		100%	52%	23%	%LL
PA4247	SeqID	103	10530	11113	11433			12221	13214	14102
,	COVERAGE	%66	%86	%66	97%		100%	%86	%86	%66
	IDENTITY	59%	52%	63%	37%		100%	48%	54%	29%
PA4248	SeqID	10344	10529	11112	11434			12220	13571	14103
	COVERAGE	100	%66	100%	%66		100%	%66	%66	100%
	IDENTITY	62%	49%	%99	20%		100%	43%	47%	62%
PA4249	Gip⇒8	10345		11111	11435				13212	14104
	COVERAGE	%66	102%	%66	100%		100%	102%	102%	%66
	IDENTITY	64%	46%	64%	40%		100%	44%	47%	64%
PA4250	SeqID	103	10599	11110					13211	14105
	COVERAGE	100%	100%	100%			100%	100%	100%	100%
	IDENTITY	69%	43%	63%			100%	46%		%19
PA4251	SeqID	10347	10527	60111	68511	11654	2915	12218	13210	14106
	COVERAGE	%66	66	%66	%66	866	100	%06	%86	%66
	IDENTITY	%69	28%	68%	48%	%69	100%	92%	%19	%89
PA4252	SeqID	10348	10526	11108					13209	14107
	COVERAGE	97%	92%	94%			100%	%86	%26	%96
	IDENTITY	65%	46%	62%			100%	46%	46%	64%
PA4253	SeqID	10349	10525	11107	11436			12216	13208	14108
	COVERAGE	101%	100%	101%	100%		100%	100%	100%	101%
	IDENTITY	85%	%99	85%			100%		%99	84%
PA4254	SeqID COVERAGE	10350 90%	10524 98%	11106	11437 84%		5170	12215 89%	13207 89%	
	IDENTITY	71%	23%				100%			

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LOCUSID Data	Data	Escherichia	ccus	Haemophilus	Helicobacter		Pseudomonas	Pseudomonas Staphylococcus Streptococcus		Salmonella
		coli	faecalis	influenzae	pylori	pneumoniae aeruginosa		aureus	oniae	typhi
PA4256	SeqID COVERAGE	10352	10560	11104	11439		5171	12260	13204	13968
	IDENTITY	%LL					100%			
PA4257	SeqID	10353	10559	11103	11592		5172	12259	13203	13969
	COVERAGE	99%	81%	100%	%66		100%	%16	93%	%66
	IDENTITY	74%	%19	72%	25%		100%	57%	29%	74%
PA4258	SeqID	10354		11102	11593		5173			13970
	COVERAGE	100	81%	100	95%		100%	%66	91%	100%
	IDENTITY	%69	21%	20%	41%		100%	48%		%69
PA4259	SeqID	10355	10557		11594			12255	13201	
	COVERAGE	100%	101%	100%	99%		100%	100%	100%	
PA4262	SeqID	10358	10549	11098	11595		5175	12240	13198	13973
	COVERAGE	100%	%56	100%	%96		100%	101%	%16	100%
	IDENTITY	%89	45%				100%	46%	44%	
PA4263		103			11442			12235	13197	13974
	COVERAGE	99%		%86	91%		100%	103%	%66	%66
	IDENTITY	75%		73%	35%		100%	46%	21%	75%
PA4264	SeqID	10360			Г	İ	5177			13975
	COVERAGE	1009	75%	100	95%	100%	100		%66	100%
	IDENTITY	%06	58%	92%	57%	92%	100%		61%	%16
PA4268	SeqID	10365			11409			12445		13967
	COVERAGE	100%	111%	100%	100%		100%	111%	111%	100%
	IDENTITY	89%	40%	89%	75%		100%			
PA4269	SeqID	10439			11410		5179	12446	13646	14042
	COVERAGE	100	100%	100%	109%		100%	101%	%66	100%
	IDENTITY	76%	46%		47%		100%	46%		75%
PA4271	SeqID	10437			11572			12449		14044
	COVERAGE	100	101%	1019	102%		100%	%86	100%	100%
	IDENTITY	%99	65%	%99	54%		100%	28%	28%	64%
PA4272	SeqID	10436	10614	11071			5181	12450		14045
	COVERAGE	% %	826	100%			100%	%66	95%	%66
	IDENTITY	%89	40%	%99			%001	39%	45%	
PA4316	SeqID	<u>ğ</u> _		11235			5182			13821
	DENTITY	88%		90%	3		100%			91%
				1			1			

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TOCOSID	Data	Escherichia coli	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella coli faecalis influenzae pylori pneumonia	Haemophilus influenzae	Helicobacter pylori	9)	r seudomonas aeruginosa	Pseudomonas Staphylococcus Streptococcus aeruginosa aureus pneumoniae		Salmonella typhi
PA4332	SeqID COVERAGE						5183 100%			
DA 4247	DENTITY					11500	100%			
7+0+1	COVERAGE					%	100%			
	IDENTITY					27%				
PA4363	SeqID	10292					5185			13742
	DENTITY	95% 40%				81%	100%			95% 41%
PA4375	SeqID	10072			11516		5186			13719
	COVERAGE	101% 33%		100%	100%		100%			101% 33%
PA4413	SeqID	10030		11188	11458			12360	13333	14077
	COVERAGE	%06	94%	92%	93%		100%	93%	%86	%06
	IDENTITY	45%	33%	41%	30%		%00I	33%		44%
PA4433	SeqID	10327		11241			5188	12237	13356	13729
	COVEKAGE	100%	%65	100%	94%	%9L 47%	100%	%55	%95%	100%
PA4473	Seath	10463		11105			5180			1-
)	COVERAGE	<u>~</u>		~			8			~
	IDENTITY	39%		37%			100%			39%
PA4506	SeqID	18801		11198			5190	12850	13248	13800
	DENTITY	58%	48%	%09 %%6	/9% 51%	91% 59%	100% 100%	46%	81% 42%	58%
PA4512	SeqID						5191			13815
	COVERAGE						100%			99%
PA4542	SeqID			Π	11489		Π	12526	13421	14088
	COVERAGE	100% 71%	101%	100%	100%		100%	101% 52%	80% 46%	100%
PA4576	SeqID						5193			
	DENTITY				_		100%			
PA4598	SeqID COVERAGE	10072 100%		11145 100%	11516 99%		5194 100%			13719 100%
	DENTILY	20%		29%	28%		100%			20%

Data Coli International Enterlocation International Colis International Colis International Colorers Internat	Critican	7 4	17.7.1		1.1	1 1111	-	1	, ,		
SeqID 10143 10826 11251 COVERAGE 100% 97% 1019 SeqID COVERAGE 100% 97% 1019 COVERAGE 10314 98% 11216 COVERAGE 107% 98% 99% DENTITY 58% 99% DENTITY 10387 99% DENTITY 88% 91% DENTITY 48% 91% SeqID 10455 91% COVERAGE 93% 91% DENTITY 43% 82% 91% SeqID 10165 90% 97% DENTITY 43% 82% 97% DENTITY 64% 82% 97% SeqID 10165 90% 11176 COVERAGE 96% 82% 97% DENTITY 64% 82% 97% SeqID 10165 90% 11176 COVERAGE 94% 82%	Jensin		coli	faecalis	raemopnuus influenzae	neucooacier pylori	a	rseuaomonas aeruginosa	r seudomonas Suepnylococcus Sireptococcus aeruginosa aureus pneumoniae		salmoneila typhi
SeqID COVERAGE DENTITY 10314 SeqID 10314 COVERAGE 107% DENTITY 58% SeqID 10387 COVERAGE 100% DENTITY 87% SeqID 10455 COVERAGE 10455 DENTITY 48% SeqID 10155 COVERAGE 93% DENTITY 43% DENTITY 90% DENTITY 90% DENTITY 43% SeqID 10165 COVERAGE 90% DENTITY 64% SeqID 10165 COVERAGE 90% DENTITY 64% SeqID 10165 COVERAGE 94% SeqID 10165 COVERAGE 94% SeqID 10167 COVERAGE 94% SeqID 11176 COVERAGE 94%		SeqID COVERAGE IDENTITY	10143 100%	10826 97% 54%	11251 101% 64%	11287 97% 52%	11675 100% 65%	5195 100% 100%	12380 98% 53%	13336 99% 50%	13979 100% 66%
SeqID		SeqID COVERAGE IDENTITY						5196 100% 100%			
SeqID 10314 11216 COVERAGE 107% 98% DENTITY 87% 99% COVERAGE 100% 99% DENTITY 87% 10972 SeqID 10455 91% COVERAGE 93% 91% DENTITY 48% 91% SeqID 10115 10619 10960 COVERAGE 86% 82% 97% DENTITY 43% 36% 97% COVERAGE 90% 11176 97% COVERAGE 90% 11176 COVERAGE 90% 11176 COVERAGE 90% 1176 COVERAGE 90% 11176 COVERAGE 94% 82% 97% COVERAGE 94% 82% 97% DENTITY 22% 33% 97% COVERAGE 94% 82% 97% DENTITY 20 33% 97% <t< td=""><td>44709</td><td>SeqID COVERAGE IDENTITY</td><td></td><td></td><td></td><td></td><td></td><td>5197 100% 100%</td><td></td><td></td><td></td></t<>	44709	SeqID COVERAGE IDENTITY						5197 100% 100%			
SeqID		SeqD COVERAGE IDENTITY	10314 107%		58%	11501 93% 39%		5198 100% 100%	12322 78% 48%	13663 91% 43%	13765 107% 58%
SeqID 10455 10972 COVERAGE 10455 10972 SeqID 48% 91% DENTITY 48% 91% SeqID 10115 10619 10960 COVERAGE 86% 82% 97% DENTITY 43% 36% 97% DENTITY 64% 82% 97% DENTITY 64% 82% 97% SeqID 10197 10796 11176 COVERAGE 94% 82% 97% DENTITY 29% 82% 97% SeqID 10197 10796 11176 COVERAGE 94% 82% 97% BENTITY 29% 82% 97% COVERAGE 94% 82% 97% DENTITY 1000 1000 1000 COVERAGE 94% 82% 97% DENTITY 1000 1000 1000 COVERAGE 1000 <t< td=""><td>4771</td><td>SeqID COVERAGE IDENTITY</td><td>10387 100%</td><td></td><td>11280 99% 75%</td><td></td><td></td><td>5199 100% 100%</td><td></td><td>13402 96% 33%</td><td>13828 97% 33%</td></t<>	4771	SeqID COVERAGE IDENTITY	10387 100%		11280 99% 75%			5199 100% 100%		13402 96% 33%	13828 97% 33%
SeqID 10455 10972 COVERAGE 93% 91% DENTITY 48% 91% SeqID 10115 10619 10960 COVERAGE 86% 82% 97% DENTITY 64% 82% 97% SeqID 10165 90% 11176 COVERAGE 94% 82% 97% DENTITY 29% 33% 97% SeqID COVERAGE 94% 82% 97% SeqID COVERAGE 94% 82% 97% DENTITY 29% 33% 97% DENTITY 20% 33% 97%	4888	SeqID COVERAGE IDENTITY						5200 100% 100%			
SeqID 10115 10619 10960 COVERAGE 86% 82% 97% DENTITY 43% 36% 97% SeqID 10165 90% 10796 11176 COVERAGE 94% 82% 97% DENTITY 29% 82% 97% SeqID 0000 10197 10796 11176 COVERAGE 94% 82% 97% SeqID COVERAGE 1000 1000 DENTITY SeqID 1000 1000		SeqID COVERAGE IDENTITY	10455 93%		10972 91% 41%			5201 100% 100%			13856 95% 48%
SeqID 10165 COVERAGE 90% DENTITY 64% SeqID 10197 10796 COVERAGE 94% 82% 97% DENTITY 29% 33% SeqID COVERAGE 1070 1070		SeqID COVERAGE IDENTITY	10115 86% 43%	10619 82% 36%	44%	11394 83% 31%		5202 100% 100%	12501 96% 37%	13458 97% 32%	14006 86% 44%
SeqID 10197 10796 11176 COVERAGE 94% 82% 97% 11176 11176 11176 11176 11176 11176 11176 11176 11177 11777		SeqID COVERAGE IDENTITY	10165 90%					5203 100% 100%			
SeqID COVERAGE IDENITIY		SeqID COVERAGE IDENTITY	10197 94% 29%	10796 82% 33%	% 27%	11383 97% 26%	11694 90% 29%	5204 100% 100%		13292 98% 30%	14057 94% 30%
	2088	SeqID COVERAGE IDENTITY						5205 100% 100%			
SeqID 10373 11126 COVERAGE 100% 96% 96% 96% 10ENTITY 41% 11%	PA5193	SeqID COVERAGE IDENTITY	10373 100%		11126 96% 39%		11709 77% 42%	5206 100% 100%			13808 100% 41%

	Salmonella typhi	13810 103% 32%		13758 89% 28%						13885 94% 52%	13748 100% 64%		
- 1										13617 94% 54%	13643 105% 39%		13236 101% 37%
	l'seudomonas Stapmytococcus Streptococcus aeruginosa aureus pneumoniae		12730 100%				12129 73% 40%			13127 94% 55%	12489 100% 38%		12623 100% 38%
	r seudomonas aeruginosa	5207 100% 100%	100	5209 100% 100%	5210 100% 100%	5211 100% 100%	5212 100% 100%	5213 100% 100%	5214 100% 100%	5215 100% 100%	5216 100% 100%	5217 100% 100%	5218 100% 100%
- 1	93	11711 52 102% 34%	1										
	Helicobacter pylori		88% 88%				11327 78% 39%			11321 94% 46%	11452 96% 35%		11609 102% 31%
**	naemopnuus influenzae		11260 100%				11158 99% 79%			11160 94% 52%	11199 100% 56%		11133 102% 58%
·	Escreticnia Enterococcus Itaemopuuns Iteucobacter Klebsteua coli faecalis influenzae pylori pneumonia	10596 71% 26%						10503 85% 28%		10924 94% 51%	10788 103% 38%		10668 102% 37%
	escnericnia coli	10375 102% 33%		10302 90% 29%			10391 100% 82%			10330 94% 52%	10413 100% 64%		10417 102% 62%
		SeqID COVERAGE IDENTITY	SeqID COVERAGE	SeqID COVERAGE IDENTITY									
TOCK	LOCUSID Data	PA5199	PA5207	PA5209	PA5248	PA5299	PA5316	PA5388			PA5443	PA5490	PA5493

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LOCUSID Data		Escherichia	Escherichia Enterococcus Haemophilus Helicobacter Klebsiella	Haemophilus	Helicobacter	Klebsiella	Pseudomonas	Pseudomonas Staphylococcus Streptococcus Salmonella	Streptococcus	Salmonella
		coli	faecalis	influenzae pylori		pneumoniae	pneumoniae aeruginosa aureus		pneumoniae typhi	typhi
PA5507 SeqID	SeqID	10119					5219			
	COVERAGE	%66					100%			
	IDENTITY	31%					100%			
PA5567	SeqID 10397		10911	69111	11450		5220	12703	13338	13923
	COVERAGE	%66	103%	%66	100%		100%	102%	101%	%66
	DENTITY	%29	39%	64%	33%		100%	34%	37%	%19

PathoSeq Cluster ID	Enterococcus faecalis	Escherichia coli	Pseudomonas aeruginosa	Staphylococcus aureus
15	EFA102326	ECO101796	PAE100280	SAU102515
55	EFA100151	ECO104157	PAE100416	SAU100633
57	EFA100617	ECO102690	PAE105434	SAU100158
1443	EFA100689	ECO103692	PAE101987	SAU100952
1861	EFA101412	ECO103231	PAE104331	SAU101793
2286	EFA103268	ECO103265	PAE104314	SAU101756
2362	EFA101425	ECO100662	PAE101537	SAU101236
2367	EFA101417	ECO103226	PAE103206	SAU101798
2549	EFA101410	ECO103233	PAE104329	SAU101791
3816	EFA101159	ECO103243	PAE104319	SAU100546
3857	EFA101415	ECO103228	PAE103204	SAU101796
4322	EFA101165	ECO103237	PAE104325	SAU100141
4569	EFA100955	ECO103217	PAE103215	SAU101808
4948	EFA101160	ECO103242	PAE104320	SAU100547
5818	EFA100742	ECO103224	PAE103208	SAU101800
8159	EFA101163	ECO103239	PAE104323	SAU100139
8296	EFA101164	ECO103238	PAE104324	SAU100140
8316	EFA101409	ECO103234	PAE104328	SAU101790
8494	EFA103062	ECO103884	PAE104311	SAU100433
8498	EFA101411	ECO103232	PAE104330	SAU101792
8499	EFA101416	ECO103227	PAE103205	SAU101797
7		ECO100071	PAE100837	SAU102674
8	EFA101340		PAE106580	SAU100118
28	EFA101403		PAE102647	SAU100514
41	EFA101753	ECO100148		SAU101565
63	EFA101685		PAE103857	SAU100331
147		ECO100645	PAE100543	SAU100053
548		ECO100377	PAE100604	SAU100747
730		ECO103592	PAE103108	SAU100061
1721	EFA101686	ECO100663		SAU101996
1749	EFA101477	ECO102557	· · · · · · · · · · · · · · · · · · ·	SAU100613
2153	EFA102656	ECO100184		SAU101869
2790	EFA102764	ECO100500		SAU101578
3164	EFA101162	ECO103240		SAU102602
3312	EFA103174		PAE105008	SAU100521
3926	EFA100194	ECO103220	***	SAU101806
4441	EFA102541		PAE105364	SAU101814
5685	EFA100190	ECO103264		SAU100157
7417	EFA102788	ECO101684		SAU102992
7437	EFA102351	ECO100084		SAU100056
7579		ECO102470	PAE102641	SAU100607
7726	EFA102551	ECO103221		SAU101805
7727	EFA100978	ECO103218		SAU101807
8092		ECO102035	PAE102964	SAU100794
8158	EFA103365		PAE104318	SAU102880
8161	EFA100210		PAE104326	SAU102527
8162	EFA101414		PAE103203	SAU101795
8092 8158 8161	EFA103365 EFA100210		PAE104318 PAE104326	SAU100794 SAU102880 SAU102527

PathoSeq	Enterococcus	Escherichia coli	Pseudomonas	Staphylococcus
Cluster ID	faecalis		aeruginosa	aureus
8164	EFA100741	ECO103223		SAU101801
8493	EFA101141		PAE104310	SAU100432
10185	EFA102728	ECO104092		SAU102578
35		ECO102870		SAU100497
44			PAE101061	SAU101143
54			PAE100225	SAU100123
85		ECO101104		SAU101262
184			PAE104901	SAU101366
362	EFA102736			SAU100414
575	EFA101790			SAU100133
579	EFA102110			SAU101624
911			PAE105432	SAU102054
941		ECO101365		SAU102162
952	EFA100615			SAU100964
1084	EFA100289	ECO102819		
1141		ECO102255		SAU102356
1232		ECO100703		SAU101346
1274			PAE103655	SAU102264
1337		ECO102562		SAU100567
1350		ECO100930	PAE103901	
1374		ECO103659		SAU101385
1427	EFA100394			SAU100714
1535		ECO101207		SAU101561
1653	EFA102655			SAU101868
1849	EFA100642			SAU101653
1932	EFA100919			SAU101365
2156	EFA101150			SAU101271
2189		ECO102827	PAE100476	
2238		ECO101436		SAU101092
2338	EFA103038			SAU100518
2411	EFA102802			SAU102246
2501	EFA101121			SAU100996
2974			PAE102537	SAU102125
3027		ECO103959		SAU200242
3239	EFA103021			SAU100300
3244	EFA100399			SAU101891
3386	EFA100426			SAU100886
3447	EFA102915			SAU102112
3460	EFA102023			SAU101399
3682	EFA100740			SAU101802
3771	EFA101540			SAU100275
4424	EFA102542	700100100	DIBIOCIO!	SAU101815
4654	TD 1 100005	ECO100488	PAE106184	0.111100550
5148	EFA100065		· · · · · · · · · · · · · · · · · · ·	SAU100658
7227	EFA100023	7001000		SAU100436
7240		ECO103672	DARIOSCO	SAU101682
7278			PAE101620	SAU301370
7374	EEA 100051		PAE106765	SAU103042
7375	EFA102051			SAU103038

PathoSeq Cluster ID	Enterococcus faecalis	Escherichia coli	Pseudomonas aeruginosa	Staphylococcus aureus
7402		ECO103572	PAE106044	<u> </u>
7419		ECO101686		SAU102693
7436	EFA101792			SAU101495
7504	EFA101670			SAU102603
7653	EFA100397			SAU100246
7660	EFA102352	ECO103698		
7719	EFA100756			SAU100496
7725	EFA100739			SAU101803
8040	EFA101736			SAU101197
8058	EFA103571			SAU101242
8077	EFA100200			SAU102231
8082	EFA101080			SAU100199
8116	EFA101963			SAU101028
8122	EFA101737			SAU101198
8141	EFA102780			SAU102433
8177	EFA103348			SAU202126
8178	EFA101022	Y)		SAU102283
8181	EFA101541			SAU102909
8191	EFA102022			SAU101398
8234	EFA103033			SAU100745
8237	EFA101682			SAU101266
8238	EFA103295			SAU100963
8251			PAE100662	SAU100596
8300	EFA101120			SAU100944
8539	EFA101339			SAU101400
8610		ECO103661		SAU102298
8874	EFA100748			SAU101155
9028	EFA103210			SAU100731
9996	EFA102338			SAU100175
10234	EFA102186			SAU102933
10248		ECO102828		SAU101220
10297	-		PAE105229	SAU101381
10328	EFA101079			SAU101547
10345	EFA100295			SAU100659
10365	EFA100641			SAU101655
10393	EFA103504			SAU100961
10402	EFA101833			SAU100880
12426	EFA101413			SAU101794
14277	EFA103081			SAU200088
14330	EFA101161			SAU102881
14455	EFA101424			SAU101771
14520	EFA100211			SAU101789
15660	EFA103375			SAU102694

EXAMPLE 13

Use of Identified Nucleic Acid Sequences as Probes

The sequences from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus 5 faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi described herein. homologous coding nucleic acids, or homologous antisense nucleic acids can be used as probes to obtain the sequence of additional genes of interest from a second cell or microorganism. For example, probes to genes encoding potential bacterial target proteins may be hybridized to nucleic acids from other organisms including other bacteria and higher organisms, to identify homologous sequences in these other organisms. For example, the identified sequences from Staphylococcus aureus, Salmonella 10 typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi, homologous coding nucleic acids, or homologous antisense nucleic acids may be used to identify homologous sequences in Anaplasma marginale, Aspergillus fumigatus, Bacillus 15 anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni. Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, 20 Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, 25 Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis and any species falling within the 30 genera of any of the above species. In some embodiments of the present invention, the nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa. Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi described herein, homologous coding nucleic acids, or homologous antisense nucleic acids may be used to identify homologous nucleic acids 35 from a heterologous organism other than E. coli.

Hybridization between the nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis,

Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi described herein, homologous coding nucleic acids, or homologous antisense nucleic acids and nucleic acids from humans might indicate that the protein encoded by the gene to which the probe corresponds is found in humans and therefore not necessarily an optimal drug target.

Alternatively, the gene can be conserved only in bacteria and therefore would be a good drug target for a broad spectrum antibiotic or antimicrobial. These probes can also be used in a known manner to isolate homologous nucleic acids from *Staphylococcus*, *Salmonella*, *Klebsiella*, *Pseudomonas*, *Enterococcus* or other cells or microorganisms, e.g. by screening a genomic or cDNA library.

Probes derived from the nucleic acid sequences from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi described herein, homologous coding nucleic acids, or homologous antisense nucleic acids, or portions thereof, can be labeled with detectable labels familiar to those skilled in the art, including radioisotopes and non-radioactive labels, to provide a detectable probe. The detectable probe can be single stranded or double stranded and can be made using techniques known in the art, including in vitro transcription, nick translation, or kinase reactions. A nucleic acid sample containing a sequence capable of hybridizing to the labeled probe is contacted with the labeled probe. If the nucleic acid in the sample is double stranded, it can be denatured prior to contacting the probe. In some applications, the nucleic acid sample can be immobilized on a surface such as a nitrocellulose or nylon membrane. The nucleic acid sample can comprise nucleic acids obtained from a variety of sources, including genomic DNA, cDNA libraries, RNA, or tissue samples.

Procedures used to detect the presence of nucleic acids capable of hybridizing to the detectable probe include well known techniques such as Southern blotting, Northern blotting, dot blotting, colony hybridization, and plaque hybridization. In some applications, the nucleic acid capable of hybridizing to the labeled probe can be cloned into vectors such as expression vectors, sequencing vectors, or in vitro transcription vectors to facilitate the characterization and expression of the hybridizing nucleic acids in the sample. For example, such techniques can be used to isolate, purify and clone sequences from a genomic library, made from a variety of bacterial species, which are capable of hybridizing to probes made from the sequences identified in Examples 5 and 6.

EXAMPLE 14

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Preparation of PCR Primers and Amplification of DNA

The identified Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi genes corresponding directly to or located within the operon of nucleic acid sequences required for proliferation, homologous coding nucleic acids, or homologous antisense nucleic acids or portions thereof can be used to prepare PCR primers for a variety of applications, including the identification or isolation of homologous sequences

from other species. For example, the Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi genes may be used to prepare PCR primers to identify or isolate homologous sequences from Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia 5 cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei. Candida kefvr (also called Candida pseudotropicalis). Candida dubliniensis. Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae, Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis or any species falling within the genera of any of the above species. In some embodiments of the present invention, the PCR primers may be used to identify or isolate homologous nucleic acids from an organism other than E. coli.

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The identified or isolated nucleic acids obtained using the PCR primers may contain part or all of the homologous nucleic acids. Because homologous nucleic acids are related but not identical in sequence, those skilled in the art will often employ degenerate sequence PCR primers. Such degenerate sequence primers are designed based on sequence regions that are either known to be conserved or suspected to be conserved such as conserved coding regions. The successful production of a PCR product using degenerate probes generated from the sequences identified herein would indicate the presence of a homologous gene sequence in the species being screened. The PCR primers are at least 10 nucleotides, and preferably at least 20 nucleotides in length. More preferably, the PCR primers are at least 20-30 nucleotides in length. In some embodiments, the PCR primers can be more than 30 nucleotides in length. It is preferred that the primer pairs have approximately the same G/C ratio, so that melting temperatures are approximately the same. A variety of PCR techniques are familiar to those skilled in the art. For a review of PCR technology, see Molecular Cloning to Genetic Engineering White, B.A. Ed. in Methods in Molecular Biology 67: Humana Press, Totowa 1997. When the entire coding sequence of the target gene is known, the 5' and 3' regions of the target gene can be used as the sequence source for PCR probe generation. In each of these PCR procedures, PCR

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primers on either side of the nucleic acid sequences to be amplified are added to a suitably prepared nucleic acid sample along with dNTPs and a thermostable polymerase such as Taq polymerase, Pfu polymerase, or Vent polymerase. The nucleic acid in the sample is denatured and the PCR primers are specifically hybridized to complementary nucleic acid sequences in the sample. The hybridized primers are extended. Thereafter, another cycle of denaturation, hybridization, and extension is initiated. The cycles are repeated multiple times to produce an amplified fragment containing the nucleic acid sequence between the primer sites.

EXAMPLE 15

Inverse PCR

The technique of inverse polymerase chain reaction can be used to extend the known nucleic acid sequence identified in Examples 5 and 6. The inverse PCR reaction is described generally by Ochman et al., in Ch. 10 of PCR Technology: Principles and Applications for DNA Amplification, (Henry A. Erlich, Ed.) W.H. Freeman and Co. (1992). Traditional PCR requires two primers that are used to prime the synthesis of complementary strands of DNA. In inverse PCR, only a core sequence need be known.

Using the sequences identified as relevant from the techniques taught in Examples 5 and 6 and applied to other species of bacteria, a subset of nucleic sequences are identified that correspond to genes or operons that are required for bacterial proliferation. In species for which a genome sequence is not known, the technique of inverse PCR provides a method for obtaining the gene in order to determine the sequence or to place the probe sequences in full context to the target sequence to which the identified nucleic acid sequence binds.

To practice this technique, the genome of the target organism is digested with an appropriate restriction enzyme so as to create fragments of nucleic acid that contain the identified sequence as well as unknown sequences that flank the identified sequence. These fragments are then circularized and become the template for the PCR reaction. PCR primers are designed in accordance with the teachings of Example 15 and directed to the ends of the identified sequence. The primers direct nucleic acid synthesis away from the known sequence and toward the unknown sequence contained within the circularized template. After the PCR reaction is complete, the resulting PCR products can be sequenced so as to extend the sequence of the identified gene past the core sequence of the identified exogenous nucleic acid sequence identified. In this manner, the full sequence of each novel gene can be identified. Additionally the sequences of adjacent coding and noncoding regions can be identified.

EXAMPLE 16

Identification of Genes Required for Escherichia coli Proliferation

Genes required for proliferation in *Escherichia coli* are identified according to the methods described above.

EXAMPLE 17

Identification of Genes Required for Neisseria gonorrhoeae Proliferation

Genes required for proliferation in *Neisseria gonorrhoeae* are identified according to the methods described above.

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EXAMPLE 18

Identification of Genes Required for Salmonella enterica Proliferation

Genes required for proliferation in Salmonella enterica are identified according to the methods described above

EXAMPLE 19

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Identification of Genes Required for Enterococcus faecium Proliferation

Genes required for proliferation in Enterococcus faecium are identified according to the

methods described above.

EXAMPLE 20

Identification of Genes Required for Haemophilus influenzae Proliferation

Genes required for proliferation in *Haemophilus influenzae* are identified according to the methods described above.

EXAMPLE 21

Identification of Genes Required for Aspergillus fumigatus Proliferation

Genes required for proliferation in *Aspergillus fumigatus* are identified according to the methods described above.

EXAMPLE 22

Identification of Genes Required for Helicobacter pylori Proliferation

Genes required for proliferation in *Helicobacter pylori* are identified according to the methods described above.

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EXAMPLE 23

Identification of Genes Required for Mycoplasma pneumoniae Proliferation

Genes required for proliferation in Mycoplasma pneumoniae are identified according to the methods described above.

EXAMPLE 24

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Identification of Genes Required for Plasmodium ovale Proliferation

Genes required for proliferation in *Plasmodium ovale* are identified according to the methods described above.

EXAMPLE 25

Identification of Genes Required for Entamoeba histolytica Proliferation

Genes required for proliferation in *Entamoeba histolytica* are identified according to the methods described above.

EXAMPLE 26

Identification of Genes Required for Candida albicans Proliferation

Genes required for proliferation in *Candida albicans* are identified according to the methods described above.

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EXAMPLE 27

Identification of Genes Required for Histoplasma capsulatum Proliferation

Genes required for proliferation in *Histoplasma capsulatum* are identified according to the methods described above.

EXAMPLE 28

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Identification of Genes Required for Salmonella typhi Proliferation

Genes required for proliferation in Salmonella typhi are identified according to the methods described above.

EXAMPLE 29

Identification of Genes Required for Salmonella paratyphi Proliferation

Genes required for proliferation in Salmonella paratyphi are identified according to the methods described above.

EXAMPLE 30

Identification of Genes Required for Salmonella cholerasuis Proliferation

Genes required for proliferation in Salmonella cholerasuis are identified according to the methods described above.

EXAMPLE 31

Identification of Genes Required for Staphylococcus epidermis Proliferation

Genes required for proliferation in *Staphylococcus epidermis* are identified according to the methods described above.

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EXAMPLE 32

Identification of Genes Required for Mycobacterium tuberculosis Proliferation

Genes required for proliferation in *Mycobacterium tuberculosis* are identified according to the methods described above.

EXAMPLE 33

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Identification of Genes Required for Mycobacterium leprae Proliferation

Genes required for proliferation in *Mycobacterium leprae* are identified according to the methods described above.

EXAMPLE 34

Identification of Genes Required for Treponema pallidum Proliferation

Genes required for proliferation in *Treponema pallidum* are identified according to the methods described above

EXAMPLE 35

Identification of Genes Required for Bacillus anthracis Proliferation

Genes required for proliferation in *Bacillus anthracis* are identified according to the methods described above.

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EXAMPLE 36

Identification of Genes Required for Yersinia pestis Proliferation

Genes required for proliferation in *Yersinia pestis* are identified according to the methods described above.

EXAMPLE 37

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Identification of Genes Required for Clostridium botulinum Proliferation

Genes required for proliferation in *Clostridium botulinum* are identified according to the methods described above.

EXAMPLE 38

Identification of Genes Required for Campylobacter jejuni Proliferation

15 Genes required for proliferation in Campylobacter jejuni are identified according to the methods described above.

EXAMPLE 39

Identification of Genes Required for Chlamydia trachomatis Proliferation

Genes required for proliferation in *Chlamydia trachomatis* are identified according to the methods described above.

EXAMPLE 40

Identification of Genes Required for Staphylococcus aureus Proliferation

Genes required for proliferation in *Staphylococcus aureus* are identified according to the methods described above.

EXAMPLE 41

Identification of Genes Required for Salmonella typhimurium Proliferation

Genes required for proliferation in Salmonella typhimurium are identified according to the methods described above.

EXAMPLE 42

Identification of Genes Required for Klebsiella Pneumoniae Proliferation

Genes required for proliferation in Klebsiella Pneumoniae are identified according to the methods described above.

EXAMPLE 43

Identification of Genes Required for Pseudomonas aeruginosa Proliferation

Genes required for proliferation in *Pseudomonas aeruginosa* are identified according to the methods described above

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EXAMPLE 44

Identification of Genes Required for Enterococcus faecalis Proliferation

Genes required for proliferation in *Enterococcus faecalis* are identified according to the methods described above.

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Use of Isolated Exogenous Nucleic Acid Fragments as Antisense Antibiotics

In addition to using the identified sequences to enable screening of molecule libraries to identify compounds useful to identify antibiotics, antisense nucleic acids complementary to the proliferation-required sequences or portions thereof, antisense nucleic acids complementary to homologous coding nucleic acids, or homologous antisense nucleic acids can be used as therapeutic agents. Specifically, the proliferation-required sequences or homologous coding nucleic acids, or portions therof, in an antisense orientation or homologous antisense nucleic acids can be provided to an individual to inhibit the translation of a bacterial target gene or the processing, folding, or assembly into a protein/RNA complex of a nontranslated RNA.

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EXAMPLE 45

Generation of Antisense Therapeutics from Identified Exogenous Sequences

Antisense nucleic acids complementary to the proliferation-required sequences described herein, or portions thereof, antisense nucleic acids complementary to homologous coding nucleic acids, or portions thereof, or homologous antisense nucleic acids or portions thereof can be used as antisense therapeutics for the treatment of bacterial infections or simply for inhibition of bacterial growth in vitro or in vivo. For example, the antisense therapeutics may be used to treat bacterial infections caused by Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi or to inhibit the growth of these organisms. The antisense therapeutics may also be used to treat infections caused by or to inhibit the growth of Anaplasma marginale, Aspergillus fumigatus, Bacillus anthracis, Bacterioides fragilis Bordetella pertussis, Burkholderia cepacia, Campylobacter jejuni, Candida albicans, Candida glabrata (also called Torulopsis glabrata), Candida tropicalis, Candida parapsilosis, Candida guilliermondii, Candida krusei, Candida kefyr (also called Candida pseudotropicalis), Candida dubliniensis, Chlamydia pneumoniae, Chlamydia trachomatus, Clostridium botulinum, Clostridium difficile, Clostridium perfringens, Coccidiodes immitis, Corynebacterium diptheriae,

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Cryptococcus neoformans, Enterobacter cloacae, Enterococcus faecalis, Enterococcus faecium, Escherichia coli, Haemophilus influenzae, Helicobacter pylori, Histoplasma capsulatum, Klebsiella pneumoniae, Listeria monocytogenes, Mycobacterium leprae, Mycobacterium tuberculosis, Neisseria gonorrhoeae, Neisseria meningitidis, Nocardia asteroides, Pasteurella haemolytica, Pasteurella multocida, Pneumocystis carinii, Proteus vulgaris, Pseudomonas aeruginosa, Salmonella bongori, Salmonella cholerasuis, Salmonella enterica, Salmonella paratyphi, Salmonella typhi, Salmonella typhimurium, Staphylococcus aureus, Listeria monocytogenes, Moxarella catarrhalis, Shigella boydii, Shigella dysenteriae, Shigella flexneri, Shigella sonnei, Staphylococcus epidermidis, Streptococcus pneumoniae, Streptococcus mutans, Treponema pallidum, Yersinia enterocolitica, Yersinia pestis or any species falling within the genera of any of the above species. In some embodiments of the present invention, the antisense therapuetics may be used to treat infection by or inhibit the growth of an organism other than E. coli.

The therapy exploits the biological process in cells where genes are transcribed into messenger RNA (mRNA) that is then translated into proteins. Antisense RNA technology contemplates the use of antisense nucleic acids, including antisense oligonucleotides, complementary to a target gene that will bind to its target nucleic acid and decrease or inhibit the expression of the target gene. For example, the antisense nucleic acid may inhibit the translation or transcription of the target nucleic acid. In one embodiment, antisense oligonucleotides can be used to treat and control a bacterial infection of a cell culture containing a population of desired cells contaminated with bacteria. In another embodiment, the antisense oligonucleotides can be used to treat an organism with a bacterial infection.

Antisense oligonucleotides can be synthesized from any of the sequences of the present invention using methods well known in the art. In a preferred embodiment, antisense oligonucleotides are synthesized using artificial means. Uhlmann & Peymann, Chemical Rev. 90:543-584 (1990) review antisense oligonucleotide technology in detail. Modified or unmodified antisense oligonucleotides can be used as therapeutic agents. Modified antisense oligonucleotides are preferred. Modification of the phosphate backbones of the antisense oligonucleotides can be achieved by substituting the internucleotide phosphate residues with methylphosphonates, phosphorothioates, phosphoramidates, and phosphate esters. Nonphosphate internucleotide analogs such as siloxane bridges, carbonate brides, thioester bridges, as well as many others known in the art may also be used. The preparation of certain antisense oligonucleotides with modified internucleotide linkages is described in U.S. Patent No. 5,142,047.

Modifications to the nucleoside units of the antisense oligonucleotides are also contemplated. These modifications can increase the half-life and increase cellular rates of uptake for the oligonucleotides *in vivo*. For example, α -anomeric nucleotide units and modified nucleotides such as 1,2-dideoxy-d-ribofuranose, 1,2-dideoxy-1-phenylribofuranose, and N^4 , N^4 -ethano-5-methyl-cytosine are contemplated for use in the present invention.

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An additional form of modified antisense molecules is found in peptide nucleic acids. Peptide nucleic acids (PNA) have been developed to hybridize to single and double stranded nucleic acids. PNA are nucleic acid analogs in which the entire deoxyribose-phosphate backbone has been exchanged with a chemically different, but structurally homologous, polyamide (peptide) backbone containing 2-aminoethyl glycine units. Unlike DNA, which is highly negatively charged, the PNA backbone is neutral. Therefore, there is much less repulsive energy between complementary strands in a PNA-DNA hybrid than in the comparable DNA-DNA hybrid, and consequently they are much more stable. PNA can hybridize to DNA in either a Watson/Crick or Hoogsteen fashion (Demidov et al., *Proc. Natl. Acad. Sci. U.S.A.* 92:2637-2641, 1995; Egholm, *Nature* 365:566-568, 1993; Nielsen et al., *Science* 254:1497-1500, 1991; Ducholm et al., *New J. Chem.* 21:19-31, 1997).

Molecules called PNA "clamps" have been synthesized which have two identical PNA sequences joined by a flexible hairpin linker containing three 8-amino-3,6-dioxaoctanoic acid units. When a PNA clamp is mixed with a complementary homopurine or homopyrimidine DNA target sequence, a PNA-DNA-PNA triplex hybrid can form which has been shown to be extremely stable (Bentin et al., *Biochemistry* 35:8863-8869, 1996; Egholm et al., *Nucleic Acids Res.* 23:217-222, 1995; Griffith et al., *J. Am. Chem. Soc.* 117:831-832, 1995).

The sequence-specific and high affinity duplex and triplex binding of PNA have been extensively described (Nielsen et al., *Science* 254:1497-1500, 1991; Egholm et al., *J. Am. Chem. Soc.* 114:9677-9678, 1992; Egholm et al., *Nature* 365:566-568, 1993; Almarsson et al., *Proc. Natl. Acad. Sci. U.S.A.* 90:9542-9546, 1993; Demidov et al., *Proc. Natl. Acad. Sci. U.S.A.* 92:2637-2641, 1995). They have also been shown to be resistant to nuclease and protease digestion (Demidov et al., *Biochem. Pharm.* 48:1010-1313, 1994). PNA has been used to inhibit gene expression (Hanvey et al., *Science* 258:1481-1485,1992; Nielsen et al., *Nucl. Acids. Res.*, 21:197-200, 1993; Nielsen et al., *Gene* 149:139-145, 1994; Good & Nielsen, Science, 95: 2073-2076, 1998), to block restriction enzyme activity (Nielsen et al., *supra.*, 1993), to act as an artificial transcription promoter (Mollegaard, *Proc. Natl. Acad. Sci. U.S.A.* 91:3892-3895, 1994) and as a pseudo restriction endonuclease (Demidov et al., *Nucl. Acids. Res.* 21:2103-2107, 1993). Recently, PNA has also been shown to have antiviral and antitumoral activity mediated through an antisense mechanism (Norton, *Nature Biotechnol.*, 14:615-619, 1996; Hirschman et al., *J. Investig. Med.* 44:347-351, 1996). PNAs have been linked to various peptides in order to promote PNA entry into cells (Basu et al., *Bioconj. Chem.* 8:481-488, 1997; Pardridge et al., *Proc. Natl. Acad. Sci. U.S.A.* 92:5592-5596, 1995).

The antisense oligonucleotides contemplated by the present invention can be administered by direct application of oligonucleotides to a target using standard techniques well known in the art. The antisense oligonucleotides can be generated within the target using a plasmid, or a phage.

Alternatively, the antisense nucleic acid may be expressed from a sequence in the chromosome of the target cell. For example, a promoter may be introduced into the chromosome of the target cell near the target gene such that the promoter directs the transcription of the antisense nucleic acid.

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Alternatively, a nucleic acid containing the antisense sequence operably linked to a promoter may be introduced into the chromosome of the target cell. It is further contemplated that the antisense oligonucleotides are incorporated in a ribozyme sequence to enable the antisense to specifically bind and cleave its target mRNA. For technical applications of ribozyme and antisense oligonucleotides see Rossi et al., **Pharmacol. Ther. 50(2):245-254**, (1991). The present invention also contemplates using a retron to introduce an antisense oligonucleotide to a cell. Retron technology is exemplified by U.S. Patent No. 5,405,775. Antisense oligonucleotides can also be delivered using liposomes or by electroporation techniques which are well known in the art.

The antisense nucleic acids described above can also be used to design antibiotic compounds comprising nucleic acids which function by intracellular triple helix formation. Triple helix oligonucleotides are used to inhibit transcription from a genome. The antisense nucleic acids can be used to inhibit cell or microorganism gene expression in individuals infected with such microorganisms or containing such cells. Traditionally, homopurine sequences were considered the most useful for triple helix strategies. However, homopyrimidine sequences can also inhibit gene expression. Such homopyrimidine oligonucleotides bind to the major groove at homopurine:homopyrimidine sequences. Thus, both types of sequences based on the sequences from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi or homologous nucleic acids that are required for proliferation are contemplated for use as antibiotic compound templates.

The antisense nucleic acids, such as antisense oligonucleotides, which are complementary to the proliferation-required nucleic acids from Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi or to homologous coding nucleic acids, or portions thereof, may be used to induce bacterial cell death or at least bacterial stasis by inhibiting target nucleic acid transcription or translation. Antisense oligonucleotides complementary to about 8 to 40 nucleotides of the proliferation-required nucleic acids described herein or homologous coding nucleic acids have sufficient complementarity to form a duplex with the target sequence under physiological conditions.

To kill bacterial cells or inhibit their growth, the antisense oligonucleotides are applied to the bacteria or to the target cells under conditions that facilitate their uptake. These conditions include sufficient incubation times of cells and oligonucleotides so that the antisense oligonucleotides are taken up by the cells. In one embodiment, an incubation period of 7-10 days is sufficient to kill bacteria in a sample. An optimum concentration of antisense oligonucleotides is selected for use.

The concentration of antisense oligonucleotides to be used can vary depending on the type of bacteria sought to be controlled, the nature of the antisense oligonucleotide to be used, and the

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relative toxicity of the antisense oligonucleotide to the desired cells in the treated culture. Antisense oligonucleotides can be introduced to cell samples at a number of different concentrations preferably between $1 \times 10^{-10} M$ to $1 \times 10^{-4} M$. Once the minimum concentration that can adequately control gene expression is identified, the optimized dose is translated into a dosage suitable for use *in vivo*. For example, an inhibiting concentration in culture of 1×10^{-7} translates into a dose of approximately 0.6 mg/kg body weight. Levels of oligonucleotide approaching 100 mg/kg body weight or higher may be possible after testing the toxicity of the oligonucleotide in laboratory animals. It is additionally contemplated that cells from the subject are removed, treated with the antisense oligonucleotide, and reintroduced into the subject. This range is merely illustrative and one of skill in the art are able to determine the optimal concentration to be used in a given case.

After the bacterial cells have been killed or controlled in a desired culture, the desired cell population may be used for other purposes.

EXAMPLE 46

Use of Antisense Oligonucleotides to Treat Contaminated Cell Cultures

The following example demonstrates the ability of an Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi antisense oligonucleotide or an antisense oligonucleotide complementary to a homologous coding nucleic acid, or portions thereof, to act as a bacteriocidal or bacteriostatic agent to treat a contaminated cell culture system. The application of the antisense oligonucleotides of the present invention are thought to inhibit the translation of bacterial gene products required for proliferation. The antisense nucleic acids may also inhibit the transcription, folding or processing of the target RNA.

In one embodiment of the present invention, the antisense oligonucleotide may comprise a phosphorothioate modified nucleic acid comprising at least about 15, at least about 20, at least about 25, at least about 30, at least about 35, at least about 40, or more than 40 consecutive nucleotides of an antisense nucleic acid listed in Table IA. A sense oligodeoxynucleotide complementary to the antisense sequence is synthesized and used as a control. The oligonucleotides are synthesized and purified according to the procedures of Matsukura, et al., Gene 72:343 (1988). The test oligonucleotides are dissolved in a small volume of autoclaved water and added to culture medium to make a 100 micromolar stock solution.

Human bone marrow cells are obtained from the peripheral blood of two patients and cultured according standard procedures well known in the art. The culture is contaminated with Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi or an organism containing a homologous nucleic acid and incubated at 37°C overnight to establish bacterial infection.

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The control and antisense oligonucleotide containing solutions are added to the contaminated cultures and monitored for bacterial growth. After a 10 hour incubation of culture and oligonucleotides, samples from the control and experimental cultures are drawn and analyzed for the translation of the target bacterial gene using standard microbiological techniques well known in the art. The target Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi gene or an organism containing the homologous coding nucleic acid is found to be translated in the control culture treated with the control oligonucleotide, however, translation of the target gene in the experimental culture treated with the antisense oligonucleotide of the present invention is not detected or reduced, indicating that the culture is no longer contaminated or is contaminated at a reduced level.

EXAMPLE 47

Use of Antisense Oligonucleotides to Treat Infections

A subject suffering from a Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi infection or an infection with an organism containing a homologous coding nucleic acid is treated with the antisense oligonucleotide preparation above. The antisense oligonucleotide is provided in a pharmaceutically acceptable carrier at a concentration effective to inhibit the transcription or translation of the target nucleic acid. The present subject is treated with a concentration of antisense oligonucleotide sufficient to achieve a blood concentration of about 0.1-100 micromolar. The patient receives daily injections of antisense oligonucleotide to maintain this concentration for a period of 1 week. At the end of the week a blood sample is drawn and analyzed for the presence or absence of the organism using standard techniques well known in the art. There is no detectable evidence of Staphylococcus aureus, Salmonella typhimurium, Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli, Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi or an organim containing a homologous coding nucleic acid and the treatment is terminated.

Antisense nucleic acids complementary to a homologous coding nucleic acid or a portion thereof may be used in the preceding method to treat individuals infected with an organism containing the homologous coding nucleic acid.

EXAMPLE 48

Preparation and Use of Triple Helix Forming Oligonucleotides

The sequences of proliferation-required nucleic acids, homologous coding nucleic acids, or homologous antisense nucleic acids are scanned to identify 10-mer to 20-mer homopyrimidine or homopyrime stretches that could be used in triple-helix based strategies for inhibiting gene expression. Following identification of candidate homopyrimidine or homopyrime stretches, their efficiency in

inhibiting gene expression is assessed by introducing varying amounts of oligonucleotides containing the candidate sequences into a population of bacterial cells that normally express the target gene. The oligonucleotides may be prepared on an oligonucleotide synthesizer or they may be purchased commercially from a company specializing in custom oligonucleotide synthesis.

The oligonucleotides can be introduced into the cells using a variety of methods known to those skilled in the art, including but not limited to calcium phosphate precipitation, DEAE-Dextran, electroporation, liposome-mediated transfection or native uptake.

Treated cells are monitored for a reduction in proliferation using techniques such as monitoring growth levels as compared to untreated cells using optical density measurements. The oligonucleotides that are effective in inhibiting gene expression in cultured cells can then be introduced *in vivo* using the techniques well known in that art at a dosage level shown to be effective.

In some embodiments, the natural (beta) anomers of the oligonucleotide units can be replaced with alpha anomers to render the oligonucleotide more resistant to nucleases. Further, an intercalating agent such as ethidium bromide, or the like, can be attached to the 3' end of the alpha oligonucleotide to stabilize the triple helix. For information on the generation of oligonucleotides suitable for triple helix formation see Griffin et al. (Science 245:967-971 (1989)).

EXAMPLE 49

Identification of Bacterial Strains from Isolated Specimens by PCR

Classical bacteriological methods for the detection of various bacterial species are time consuming and costly. These methods include growing the bacteria isolated from a subject in specialized medium, cultivation on selective agar medium, followed by a set of confirmation assays that can take from 8 to 10 days or longer to complete. Use of the identified sequences of the present invention provides a method to dramatically reduce the time necessary to detect and identify specific bacterial species present in a sample.

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In one exemplary method, bacteria are grown in enriched medium and DNA samples are isolated from specimens of, for example, blood, urine, stool, saliva or central nervous system fluid by conventional methods. A panel of PCR primers based on identified sequences unique to various species or types of cells or microorganisms are then utilized in accordance with Example 12 to amplify DNA of approximately 100-200 nucleotides in length from the specimen. A separate PCR reaction is set up for each pair of PCR primers and after the PCR reaction is complete, the reaction mixtures are assayed for the presence of PCR product. The presence or absence of bacteria from the species to which the PCR primer pairs belong is determined by the presence or absence of a PCR product in the various test PCR reaction tubes.

Although the PCR reaction is used to assay the isolated sample for the presence of various bacterial species, other assays such as the Southern blot hybridization are also contemplated.

Compounds which inhibit the activity or reduce the amount of gene products required for proliferation may be identified using rational drug design. These methods may be used with the

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proliferation-required polypeptides described herein or homologous polypeptides. In such methods, the structure of the gene product is determined using methods such as x-ray crystallography, NMR, or computer modelling. Compounds are screened to identify those which have a structure which allows them to interact with the gene product. In some embodiments, the compounds are screened to identify those which have structures which allow them to interact with regions of the gene product which are important for its activity. For example, the compounds may be screened to identify those which have structures which allow them to bind to the active site of the gene product to inhibit its activity. For example, the compound may be a suicide substrate which binds to the active site with high affinity, thereby preventing the gene product from acting on its natural substrate. Alternatively, the compound may bind to a region of the gene product which is involved in complex formation with other biomolecules. In such instances, the activity of the gene product is inhibited by blocking the interaction between the gene product and other members of the complex.

Thus, one embodiment of the present invention comprises a method of using a crystal of the gene products of the present invention and/or a dataset comprising the three-dimensional coordinates obtained from the crystal in a drug-screening assay. The present invention also includes agents (modulators or drugs) that are identified by the methods of the present invention, along with the method of using agents (modulators or drugs) identified by a method of the present invention, for inhibiting the activity of or modulating the amount of an essential gene product. The present invention also includes crystals comprising the gene products of the present invention or portions thereof.

In some embodiments of the present invention, the three-dimensional structure of the polypeptides required for proliferation is determined using X-ray crystallography or NMR. The coordinates of the determined structure are used in computer-assisted modeling programs to identify compounds that bind to and/or modulate the activity or amount of the encoded polypeptide. The method may include the following steps: 1) the generation of high-purity crystals of the encoded recombinant (or endogenous) polypeptide for analysis; 2) determination of the three-dimensional structure of the polypeptide; and, 3) the use of computer-assisted "docking" programs to analyze the molecular interaction of compound structure and the polypeptide (i.e., drug screening).

General methods for performing each of the above steps are described below and are also well known to those of skill in the art. Any method known to those of skill in the art, including those described herein, may be employed for generating the three-dimensional structure for each identified essential gene product and its use in the drug-screening assays.

Crystals of the gene products required for proliferation may be obtained as follows. Under certain conditions, molecules condense from solution into a highly-ordered crystalline lattice, which is defined by a unit cell, the smallest repeating volume of the crystalline array. The contents of such a cell can interact with and diffract certain electromagnetic and particle waves (e.g., X-rays,

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neutron beams, electron beams etc.). Due to the symmetry of the lattice, the diffracted waves interact to create a diffraction pattern. By measuring the diffraction pattern, crystallographers are able to reconstruct the three-dimensional structure of the atoms in the crystal.

Any method known to those of skill in the art, including those set forth below, may be employed to prepare high-purity crystals. For example, crystals of the product of the identified essential gene can be grown by a number of techniques including batch crystallization, vapor diffusion (either by sitting drop or hanging drop) and by microdialysis. Seeding of the crystals in some instances is required to obtain X-ray quality crystals. Standard micro and/or macro seeding of crystals may therefore be used. Exemplified below is the hanging-drop vapor diffusion procedure. Hanging drops of an essential gene product (2.5 µl, 10 mg/ml) in 20 mM Tris, pH=8.0, 100 mM NaCl are mixed with an equal amount of reservoir buffer containing 2.7-3.2 M sodium formate and 100 mM Tris buffer, pH=8.0, and kept at 4°C. Crystal showers may appear after 1-2 days with large single crystals growing to full size (0.3 X 0.3 X 0.15 mm³) within 2-3 weeks. Crystals are harvested in 3.5 M sodium formate and 100 mM Tris buffer, pH=8.0 and cryoprotected in 3.5 M sodium formate, 100 mM Tris buffer, pH=8.0, 10% (w/v) sucrose, and 10% (v/v) ethylene glycol before flash freezing in liquid propane. In some embodiments, the crystal may be obtained using the methods described in U.S. Patent No. 5,869,604. The method involves (a) contacting a mixture containing uncrystallized polypeptides with an exogenous nucleating agent that has an areal lattice match of at least 90.4% to the polypeptide,(b) crystallizing the polypeptides, thereby forming at least one crystal of the polypeptide attached to the nucleating agent, the attached crystal being of a high purity, and at least one polypeptide crystal unattached to the nucleating agent, the unattached crystal being of a lower purity than the attached crystal, and (c) separating the crystal attached to the nucleating agent from the crystal unattached to the nucleating agent. The crystallized polypeptide may also be purified from contaminants by (a) contacting a mixture containing uncrystallized polypeptides and a contaminant with an exogenous nucleating agent that has an areal lattice match of at least 90.4% to the polypeptide, (b) crystallizing the polypeptides, thereby forming at least one crystal of the polypeptide attached to the nucleating agent, the attached crystal being of a high purity and produced in a high yield, and at least one crystal unattached to the nucleating agent, the unattached crystal being of a lower purity than the attached crystal, and (c) separating the crystal attached to the nucleating agent from the crystal unattached to the nucleating agent.

Once a crystal of the present invention is grown, X-ray diffraction data can be collected using methods familiar to those skilled in the art. Therefore, any person with skill in the art of protein crystallization having the present teachings and without undue experimentation can crystallize a large number of alternative forms of the essential gene products from a variety of different organisms, or polypeptides having conservative substitutions in their amino acid sequence.

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A crystal lattice is defined by the symmetry of its unit cell and any structural motifs the unit cell contains. For example, there are 230 possible symmetry groups for an arbitrary crystal lattice, while the unit cell of the crystal lattice group may have an arbitrary dimension that depends on the molecules making up the lattice. Biological macromolecules, however, have asymmetric centers and are limited to 65 of the 230 symmetry groups. See Cantor et al., Biophysical Chemistry, Vol. III, W. H. Freeman & Company (1980).

A crystal lattice interacts with electromagnetic or particle waves, such as X-rays or electron beams respectively, that have a wavelength with the same order of magnitude as the spacing between atoms in the unit cell. The diffracted waves are measured as an array of spots on a detection surface positioned adjacent to the crystal. Each spot has a three-dimensional position, hkl, and an intensity, I(hkl), both of which are used to reconstruct the three-dimensional electron density of the crystal with the so-called Electron Density Equation. The Electron Density Equation states that the three-dimensional electron density of the unit cell is the Fourier transform of the structure factors. Thus, in theory, if the structure factors are known for a sufficient number of spots in the detection space, then the three-dimensional electron density of the unit cell could be calculated using the Electron Density Equation.

In some embodiments of the present invention, an image of a crystal of a gene product required for proliferation or a portion thereof is obtained with the aid of a digital computer and the crystal's diffraction pattern as described in U.S. Patent No. 5,353,236. The diffraction pattern contains a plurality of reflections, each having an associated resolution. The image is obtained by (a) converting the diffraction pattern of the crystal into computer usable normalized amplitudes, the pattern being produced with a diffractometer; (b) determining from the diffraction pattern a dimension of a unit cell of the crystal; (c) providing an envelope defining the region of the unit cell occupied by the gene product or portion thereof in the crystal; (d) distributing a collection of scattering bodies within said envelope, the collection of scattering bodies having various arrangements, each of which has an associated pattern of Fourier amplitudes; (e) condensing the collection of scattering bodies to a condensed arrangement that results in a high correlation between a diffraction pattern and the pattern of Fourier amplitudes for said collection of scattering bodies; (f) determining the phase associated with at least one of the reflections of said diffraction pattern from the condensed arrangement of scattering bodies; (g) calculating an electron density distribution of the gene product or portion thereof within the unit cell from the phase determined in procedure f; and (h) displaying a graphical image of the gene product or portion thereof constructed from said electron density distribution.

The crystals of the gene products required for proliferation may be used in drug screening methods such as those described in U.S. Patent Number 6,156,526. Briefly, in such methods, a compound which inhibits the formation of a complex comprising the gene product or a portion thereof is identified as follows. A set of atomic coordinates defining the three-dimensional

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structure of a complex including the gene product of interest or a portion thereof are determined. A potential compound that binds to the gene product or a portion thereof involved in complex formation is selected using the atomic coordinates obtained above. The compound is contacted with the gene product or portion thereof and its binding partner(s) in the complex under conditions which would permit the complex to form in the absence of the potential compound. The binding affinity of the gene product or portion thereof for its binding partner(s) is determined and a potential compound is identified as a compound that inhibits the formation of the complex when there is a decrease in the binding affinity of the gene product or portion thereof for its binding partner(s).

In some embodiments of the present invention, the three dimensional structure of the essential gene product is determined and potential agonists and/or potential antagonists are designed with the aid of computer modeling [Bugg et al., Scientific American, Dec.:92-98 (1993); West et al., TIPS, 16:67-74 (1995); Dunbrack et al., Folding & Design, 2:27-42 (1997)].

Computer analysis may be performed with one or more of the computer programs including: QUANTA, CHARMM, INSIGHT, SYBYL, MACROMODEL and ICM [Dunbrack et al., Folding & Design, 2:27-42 (1997)]. In a further embodiment of this aspect of the invention, an initial drug-screening assay is performed using the three-dimensional structure so obtained, preferably along with a docking computer program. Such computer modeling can be performed with one or more Docking programs such as FlexX, DOC, GRAM and AUTO DOCK [Dunbrack et al., Folding & Design, 2:27-42 (1997)].

It should be understood that for each drug screening assay provided herein, a number of iterative cycles of any or all of the steps may be performed to optimize the selection. The drug screening assays of the present invention may use any of a number of means for determining the interaction between an agent or drug and an essential gene product.

In some embodiments of the present invention, a drug can be specifically designed to bind to an essential gene product of the present invention through NMR based methodology. [Shuker et al., pi Science 274:1531-1534 (1996).] NMR spectra may be recorded using devices familiar to those skilled in the art, such as the Varian Unity Plus 500 and unity 600 spectrometers, each equipped with a pulsed-field gradient triple resonance probe as analyzed as described in Bagby et al., [Cell 82:857-867 (1995)]. Sequential resonance assignments of backbone ¹H, .¹⁵ N, and .¹³ C atoms may be made using a combination of triple resonance experiments similar to those previously described [Bagby et al., Biochemistry, 33:2409-2421 (1994a)], except with enhanced sensitivity [Muhandiram and Kay, J. Magn. Reson., 103: 203-216 (1994)] and minimal H₂O saturation [Kay et al., J. Magn. Reson., 109:129-133 (1994)]. Side chain ¹H and ¹³ C assignments may be made using HCCH-TOCSY [Bax et al., J. Magn. Reson., 87:620-627 (1990)] experiments with mixing times of 8 ms and 16 ms.in solution but need not be included in structure calculations. Nuclear Overhauser effect (NOE) cross peaks in two-dimensional ¹H--¹H NOE spectroscopy (NOESY), three-dimensional ¹⁵N-edited NOESY-HSQC [Zhang et al., J. Biomol, NMR, 4:845-858 (1994)] and

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three-dimensional simultaneous acquisition ¹⁵ N/¹³C-edited NOE [Pascal et al., J. Magn. Reson., 103:197-201 (1994)] spectra may be obtained with 100 ms NOE mixing times. Standard pseudo-atom distance corrections [Wuthrich et al., J. Mol. Biol., 169:949-961 (1983)] may be incorporated to account for center averaging. An additional 0.5 .ANG. may be added to the upper limits for distances involving methyl groups [Wagner et al., J. Mol. Biol., 196:611-639 (1987); Clore et al., Biochemistry, 26:8012-8023 (1987)].

The structures can be calculated using a simulated annealing protocol [Nilges et al., In computational Aspects of the Study of Biological Macromolecules by Nuclear Magnetic Resonance Spectroscopy, J. C. Hoch, F. M. Poulsen, and C. Redfield, eds., New York: Plenum Press, pp. 451-455 (1991)] within X-PLOR [Brunger, X-PLOR Manual, Version 3.1, New Haven, Conn.: Department of Molecular Biophysics and Biochemistry, Yale University (1993)] using the previously described strategy [Bagby et al., Structure, 2:107-122 (1994b)]. Interhelical anges may be calculated using a program written by K. Yap. Accessible surface areas were calculated using the program Naccess, available from Prof. J. Thornton, University College, London.

Compounds capable of reducing the activity or amount of gene products required for cellular proliferation may be identified using the methods described in US Pat. No. 6.077.682. Briefly, the three-dimensional structure of the gene product or portion thereof may be used in a drug screening assay by (a) selecting a potential drug by performing rational drug design with the three-dimensional structure determined from one or more sets of atomic coordinates of the gene product or portion thereof in conjunction with computer modeling; (b) contacting the potential drug with a polypeptide comprising the gene product or portion thereof and (c) detecting the binding of the potential drug with said polypeptide; wherein a potential drug is selected as a drug if the potential drug binds to the polypeptide. In some methods, the three-dimensional structure of the gene product or portion thereof is used in a drug screening assay involving (a) selecting a potential drug by performing structural based rotational drug design with the three-dimensional structure of the gene product or portion thereof; wherein said selecting is performed in conjunction with computer modeling; (b) contacting the potential drug with a polypeptide comprising the gene product or portion thereof in the presence of a substrate of the gene product; wherein in the absence of the potential drug the substrate is acted upon by the gene product; and (c) determining the extent to which the gene product acted upon the substrate; wherein a drug is selected when a decrease in the action of the gene product on the substrate is determined in the presence of the potential drug relative to in its absence. In some embodiments, the preceding method further involves(d) contacting the potential drug with the gene product or portion thereof for NMR analysis; wherein a binding complex forms between the potential drug and said gene product or portion thereof for NMR analysis; wherein the gene product or portion thereof for NMR analysis comprises a conservative amino acid substitution; (e) determining the three-dimensional structure of the binding complex by NMR; and (f) selecting a candidate drug by performing structural based rational drug

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design with the three-dimensional structure determined for the binding complex; wherein said selecting is performed in conjunction with computer modeling; (g) contacting the candidate drug with a second polypeptide comprising the gene product or portion thereof in the presence of a substrate of the gene product or portion thereof; wherein in the absence of the candidate drug the substrate is acted upon by the second polypeptide; and (h) determining the amount of action of the second polypeptide on the substrate; wherein a drug is selected when a decrease in the amount of action of the second polypeptide is determined in the presence of the candidate drug relative to in its absence.

Once the three-dimensional structure of a crystal comprising an essential gene product is determined, a potential modulator of its activity, can be examined through the use of computer modeling using a docking program such as FlexX, GRAM, DOCK, or AUTODOCK [Dunbrack et al., 1997, supra], to identify potential modulators. This procedure can include computer fitting of potential modulators to the polypeptide or fragments thereof to ascertain how well the shape and the chemical structure of the potential modulator will bind. Computer programs can also be employed to estimate the attraction, repulsion, and steric hindrance of the two binding partners (e.g., the essential gene product and a potential modulator). Generally the tighter the fit, the lower the steric hindrances, and the greater the attractive forces, the more potent the potential modulator since these properties are consistent with a tighter binding constant. Furthermore, the more specificity in the design of a potential drug the more likely that the drug will not interact as well with other proteins. This will minimize potential side-effects due to unwanted interactions with other proteins.

Compound and compound analogs can be systematically modified by computer modeling programs until one or more promising potential analogs is identified. In addition systematic modification of selected analogs can then be systematically modified by computer modeling programs until one or more potential analogs are identified. Such analysis has been shown to be effective in the development of HIV protease inhibitors [Lam et al., Science 263:380-384 (1994); Wlodawer et al., Ann. Rev. Biochem. 62:543-585 (1993); Appelt, Perspectives in Drug Discovery and Design 1:23-48 (1993); Erickson, Perspectives in Drug Discovery and Design 1:109-128 (1993)]. Alternatively a potential modulator could be obtained by initially screening a random peptide library produced by recombinant bacteriophage for example, [Scott and Smith, Science, 249:386-390 (1990); Cwirla et al., Proc. Natl. Acad. Sci., 87:6378-6382 (1990); Devlin et al., Science, 249:404-406 (1990)]. A peptide selected in this manner would then be systematically modified by computer modeling programs as described above, and then treated analogously to a structural analog.

Example 45 describes computer modelling of the structures of gene products required for proliferation.

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EXAMPLE 50

Determination of the Structure of Gene Products Required for Proliferation Using Computer Modelling

Three dimensional models were built by applying computer modelling methods to some of the gene products required for proliferation of *Staphylococcus aureus* using the amino acid sequences of the encoded proteins as follows. Sir Tom Blundell's program COMPOSER as provided by Tripos Associates in their BIOPOLYMER module to SYBYL was used to build the models. Skolnik's method of topology fingerprinting as implemented in Matchmaker was used to score the average mutation free energy. This number is in Boltzmans (units of kT) and should be negative (the more negative, the better the model.

Composer uses a Needleman Wunsch alignment with jumbling to find significant alignments. The reported parameters are percent identity and significance as measured from the jumbling. Those matches which were 30% identical and had a significance greater that 4 on the scale were judged to be good candidates for model building templates. If no three dimensional structures met these criteria, then a BLAST search was conducted against the most recent PDB sequence database. Any significant hits discovered in this manner were then added to the binary protein structure database and the candidate search was repeated in the manner discussed above.

In the next phase, Composer assigned structurally conserved and structurally variable regions and built the backbone structure and then searched the database for structures of the variable loops. These were then spliced in and a model of the protein resulted. Any loops (variable regions) which were unassignable were manually built and refined with a combination of dynamics.

The structure was then refined. Hydrogen atoms were added and a non-active aggregate was defined. 1000pS of dynamics using AMBER ALL-ATOM and Kollman charges are performed. Next a minimization cycle of up 5000 steepest decent steps were performed and then the aggregate was thawed and the process was repeated on the entire protein.

The resulting structure was then validated in MATCHMAKER. The topologically scanned free energy determined from empirically derived protein topologies was computed and the average energy/residue is reported in Boltzamans was reported. As this number represents a free energy the more negative it is the more favorable it is.

Sixty six proteins required for the proliferation of *Staphylococcus aureus* were modelled as described above. MATCHMAKER energies were computed for these. The distribution of the models built by class is shown in the table below.

WO 01/70955 PCT/US01/09180

Classification	Number of Models	Average Matchmaker Energy
Acylases	1	-0.10
Dehydrogenases	3	-0.12
DNA Related	3	-0.12
Heat Shock Protein	2	-0.16
Hydrolases	3	-0.16
Isomerases	1	0.05
Ligases	7	-0.07
Lyases	1	-0.09
Membrane Anchored	1	-0.12
Misc	18	-0.21
Oxidoreductases	6	-0.09
Proteases	1	-0.03
Ribosome	3	-0.11
Synthases	4	-0.14
Transferases	6	-0.12

Table 1. Distribution of models built with their MATCHMAKER energies in kT

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The validity of the above method was confirmed using FtsZ. In the case of FtsZ, a crystal structure from M. Janeschi was available. Examination of the gross structural features determined using the above modelling showed all of the folds in the correct place, although there were some minor differences from the structure determined by x-ray crystallography.

EXAMPLE 51

FUNCTIONAL COMPLEMENTATION

10 In another embodiment, gene products whose activities may be complemented by a proliferation-required gene product from Staphylococcus aureus, Salmonella typhimurium. Klebsiella pneumoniae, Pseudomonas aeruginosa, Enterococcus faecalis, Escherichia coli. Enterococcus faecalis, Haemophilus influenzae, Helicobacter pylori, or Salmonella typhi or homologous polypeptides are identified using merodiploids, created by introducing a plasmid or 15 Bacterial Artificial Chromosome into an organism having a mutation in the essential gene which reduces or eliminates the activity of the gene product. In some embodiments, the mutation may be a conditional mutation, such as a temperature sensitive mutation, such that the organism proliferates under permissive conditions but is unable to proliferate under non-permissive conditions in the absence of complementation by the gene on the plasmid or Bacterial Artificial Chromosome. 20 Alternatively, duplications may be constructed as described in Roth et al. (1987) Biosynthesis of Aromatic Amino Acids in Escherichia coli and Salmonella typhimurium, F. C. Neidhardt, ed., American Society for Microbiology, publisher, pp. 2269-2270. Such methods are familiar to those skilled in the art.

Table VIII provides a cross reference for SEQ ID NOs. of the nucleotide sequences discussed herein and the SEQ ID NOs. of the polypeptides encoded by these nucleotide.

Nucleotide SeqID	Protein SeqID
5916	10013
5917	10014
5918	10015
5919	10016
5920	10017
5921	10018
5922	10019
5923	10020
5924	10021
5925	10022
5926	10023
5927	10024
5928	10025
5929	10026
5930	10027
5931	10027
5932	10029
5933	10029
5934	10030
5935	10031
5936	10032
5937	10033
5938	10034
5939	10033
5940	10037
5941	10037
5942	10038
5942	10039
5943	10040
5945	10041
5946	10042
5947	10044
5948	10045
5949	10046
5950	10047
5951	10048
5952	10049
5953	10050
5954	10051
5955	10052
5956	10053
5957	10054
5958	10055
5959	10056
5960	10057
5961	10058
5962	10059

Nucleotide SeqID	Protein SeqID
5963	10060
5964	10061
5965	10062
5966	10063
5967	10064
5968	10065
5969	10066
5970	10067
5971	10068
5972	10069
5973	10070
5974	10071
5975	10072
5976	10073
5977	10074
5978	10075
5979	10076
5980	10077
5981	10078
5982	10079
5983	10080
5984	10081
5985	10082
5986	10083
5987	10084
5988	10085
5989	10086
5990	10087
5991	10088
5992	10089
5993	10090
5994	10091
5995	10092
5996	10093
5997	10094
5998	10095
5999	10096
6000	10097
6001	10098
6002	10099
6003	10100
6004	10101
6005	10102
6006	10103
6007	10104
6008	10105
6009	10106

Nucleotide SeqID	Protein SeqID
6010	10107
6011	10108
6012	10109
6013	10110
6014	10111
6015	10112
6016	10113
6017	10114
6018	10115
6019	10116
6020	10117
6021	10118
6022	10119
6023	10120
6024	10121
6025	10122
6026	10123
6027	10124
6028	10125
6029	10126
6030	10127
6031	10128
6032	10129
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6036	10133
6037	10134
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6040	10137
6041	10138
6042	10139
6043	10140
6044	10141
6045	10142
6046	10143
6047	10144
6048	10145
6049	10146
6050	10147
6051	10148
6052	10149
6053	10150
6054	10151
6055	10152
6056	10153
6057	10154

Nucleotide SeqID	Protein SeqID
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6058	10155
6059	10156
6060	10157
6061	10158
6062	10159
6063	10160
6064	10161
6065	10162
6066	10163
6067	10164
6068	10165
6069	10166
6070	10167
6071	10168
6072	10169
6073	10170
6074	10171
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6086	10183
6087	10184
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6089	10186
6090	10187
6091	10188
6092	10189
6093	10190
6094	10191
6095	10192
6096	10193
6097	10194
6098	10195
6099	10196
6100	10197
6101	10198
6102	10199
6103	10200
6104	10201
6105	10202

Nucleotide SeqID	Protein SeqID
6106	10203
6107	10204
6108	10205
6109	10206
6110	10207
6111	10208
6112	10209
6113	10210
6114	10211
6115	10212
6116	10213
6117	10214
6118	10215
6119	10216
6120	10217
6121	10218
6122	10219
6123	10220
6124	10221
6125	10222
6126	10223
6127	10224
6128	10225
6129	10226
6130	10227
6131	10228
6132	10229
6133	10230
6134	10231
6135	10232
6136	10233
6137	10234
6138	10235
6139	10236
6140	10237
6141	10238
6142	10239
6143	10240
6144	10241
6145	10242
6146	10243
6147	10244
6148	10245
6149	10246
6150	10247
6151	10248
6152	10249
6153	10250

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9951	14049
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9953	14051
9954	14052
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9961	14059
9962	14060
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9965	14063
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9967	14065
9968	14066
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Nucleotide SeqID	Protein SeqID
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9969	14067
9970	14068
9971	14069
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9973	14071
9974	14072
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9978	14076
9979	14077
9980	14078
9981	14079
9982	14080
9983	14081
9984	14082
9985	14083
9986	14084
9987	14085
9988	14086
9989	14087
9990	14088
9991	14089
9992	14090
9993	14091
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9998	14096
9999	14097
10000	14098
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10006	14104
10007	14105
10008	14106
10009	14107
10010	14108
10011	14109
10012	14110

SeqID	Clone name	Organism
8	E3M10000001A02	Enterococcus faecalis
9	E3M10000001A06	Enterococcus faecalis
10	E3M10000001B01	Enterococcus faecalis
11	E3M10000001B02	Enterococcus faecalis
12	E3M10000001B05	Enterococcus faecalis
13	E3M10000001B06	Enterococcus faecalis
14	E3M10000001B08	Enterococcus faecalis
15	E3M10000001B10	Enterococcus faecalis
16	E3M10000001C02	Enterococcus faecalis
17	E3M10000001C09	Enterococcus faecalis
18	E3M10000001D02	Enterococcus faecalis
19	E3M10000001D04	Enterococcus faecalis
20	E3M10000001D05	Enterococcus faecalis
21	E3M10000001D09	Enterococcus faecalis
22	E3M10000001E01	Enterococcus faecalis
23	E3M10000001E02	Enterococcus faecalis
24	E3M10000001E03	Enterococcus faecalis
25	E3M10000001E04	Enterococcus faecalis
26	E3M10000001E08	Enterococcus faecalis
27	E3M10000001E09	Enterococcus faecalis
28	E3M10000001F02	Enterococcus faecalis
29	E3M10000001F04	Enterococcus faecalis
30	E3M10000001F06	Enterococcus faecalis
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33	E3M10000001G03	Enterococcus faecalis
34	E3M10000001G04	Enterococcus faecalis
35	E3M1000001G05	Enterococcus faecalis
36	E3M10000001H02	Enterococcus faecalis
37	E3M10000001H03	Enterococcus faecalis
38	E3M10000001H04	Enterococcus faecalis
39	E3M10000004A04	Enterococcus faecalis
40	E3M1000004C03	Enterococcus faecalis
41	E3M1000004D01	Enterococcus faecalis
42	E3M10000004D02	Enterococcus faecalis
43	E3M1000004D10	Enterococcus faecalis
44	E3M10000004E11	Enterococcus faecalis
45	E3M10000004F08	Enterococcus faecalis
46	E3M10000004F10	Enterococcus faecalis
47	E3M1000004G01	Enterococcus faecalis
48	E3M10000004H11	Enterococcus faecalis
49	E3M1000005A07	Enterococcus faecalis
50	E3M1000005B01	Enterococcus faecalis
51	E3M1000005B08	Enterococcus faecalis
52	E3M1000005C01	Enterococcus faecalis
53	E3M1000005C03	Enterococcus faecalis
54	E3M1000005C04	Enterococcus faecalis
55	E3M1000005D03	Enterococcus faecalis

WO 01/70955 TABLE IA PCT/US01/09180

SeqID	Clone name	Organism
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58	E3M10000005E01	Enterococcus faecalis
59	E3M10000005E02	Enterococcus faecalis
60	E3M10000005E03	Enterococcus faecalis
61	E3M10000005E08	Enterococcus faecalis
62	E3M10000005F07	Enterococcus faecalis
63	E3M10000005F10	Enterococcus faecalis
64	E3M10000005G05	Enterococcus faecalis
65	E3M10000005H04	Enterococcus faecalis
66	E3M10000006B03	Enterococcus faecalis
67	E3M10000006C01	Enterococcus faecalis
68	E3M10000006C12	Enterococcus faecalis
69	E3M10000006D03	Enterococcus faecalis
70	E3M10000006E11	Enterococcus faecalis
71	E3M10000006F04	Enterococcus faecalis
72	E3M10000006G04	Enterococcus faecalis
73	E3M10000006G12	Enterococcus faecalis
74	E3M10000006H09	Enterococcus faecalis
75	E3M10000007A02	Enterococcus faecalis
76	E3M1000007B02	Enterococcus faecalis
77	E3M1000007B03	Enterococcus faecalis
78	E3M10000007C03	Enterococcus faecalis
79	E3M1000007C04	Enterococcus faecalis
80	E3M1000007D03	Enterococcus faecalis
81	E3M10000007E05 E3M10000007F01	Enterococcus faecalis
83	E3M1000007F01	Enterococcus faecalis Enterococcus faecalis
84	E3M1000007F06	Enterococcus faecalis Enterococcus faecalis
85	E3M10000007G01	Enterococcus faecalis Enterococcus faecalis
86	E3M1000008C08	Enterococcus faecalis Enterococcus faecalis
87	E3M1000008C09	Enterococcus faecalis
88	E3M1000008D08	Enterococcus faecalis
89	E3M10000008E02	Enterococcus faecalis
90	E3M10000008G05	Enterococcus faecalis
91	E3M1000008G09	Enterococcus faecalis
92	E3M10000008H02	Enterococcus faecalis
93	E3M1000009C07	Enterococcus faecalis
94	E3M1000009C09	Enterococcus faecalis
95	E3M1000009D01	Enterococcus faecalis
96	E3M1000009E02	Enterococcus faecalis
97	E3M1000009E03	Enterococcus faecalis
98	E3M1000009E05	Enterococcus faecalis
99	E3M10000009G02	Enterococcus faecalis
100	E3M10000010C08	Enterococcus faecalis
101	E3M10000010D05	Enterococcus faecalis
102	E3M10000010F01	Enterococcus faecalis
103	E3M10000010G05	Enterococcus faecalis
104	E3M10000010G07	Enterococcus faecalis

SeqID	Clone name	Organism
105	E3M10000010G09	Enterococcus faecalis
106	E3M10000010G10	Enterococcus faecalis
107	E3M10000010H02	Enterococcus faecalis
108	E3M10000011A09	Enterococcus faecalis
109	E3M10000011B03	Enterococcus faecalis
110	E3M10000011B09	Enterococcus faecalis
111	E3M10000011C07	Enterococcus faecalis
112	E3M10000011D03	Enterococcus faecalis
113	E3M10000011H02	Enterococcus faecalis
114	E3M10000011H05	Enterococcus faecalis
115	E3M10000012B01	Enterococcus faecalis
116	E3M10000012B02	Enterococcus faecalis
117	E3M10000012B07	Enterococcus faecalis
118	E3M10000012B08	Enterococcus faecalis
119	E3M10000012C01	Enterococcus faecalis
120	E3M10000012D10	Enterococcus faecalis
121	E3M10000012E08	Enterococcus faecalis
122	E3M10000012F05	Enterococcus faecalis
123	E3M10000012F06	Enterococcus faecalis
124	E3M10000012F07	Enterococcus faecalis
125	E3M10000012F10	Enterococcus faecalis
126	E3M10000012G02	Enterococcus faecalis
127	E3M10000012G07	Enterococcus faecalis
128	E3M10000013A06	Enterococcus faecalis
129	E3M10000013A07	Enterococcus faecalis
130	E3M10000013C05	Enterococcus faecalis
131	E3M10000013D02	Enterococcus faecalis
132	E3M10000013D08	Enterococcus faecalis
133	E3M10000013D10	Enterococcus faecalis
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136	E3M10000013F05	Enterococcus faecalis
137	E3M10000013F12	Enterococcus faecalis
138	E3M10000013G10	Enterococcus faecalis
139	E3M10000013H03	Enterococcus faecalis
140	E3M10000013H05	Enterococcus faecalis
141	E3M10000013H10	Enterococcus faecalis
142	E3M10000014B12	Enterococcus faecalis
143	E3M10000014E12	Enterococcus faecalis
144	E3M10000014G09	Enterococcus faecalis
145	E3M10000015B04	Enterococcus faecalis
146	E3M10000015B12	Enterococcus faecalis
147	E3M10000015E12	Enterococcus faecalis
148	E3M10000016A03	Enterococcus faecalis
149	E3M10000016A04	Enterococcus faecalis
150	E3M10000016C11	Enterococcus faecalis
151	E3M10000016D03	Enterococcus faecalis
152	E3M10000016F06	Enterococcus faecalis
153	E3M10000016F10	Enterococcus faecalis

TABLE IA

SeqID	Clone name	Organism
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156	E3M10000017A09	Enterococcus faecalis
157	E3M10000017D09	Enterococcus faecalis
158	E3M10000018A07	Enterococcus faecalis
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162	E3M10000018H06	Enterococcus faecalis
163	E3M10000019B06	Enterococcus faecalis
164	E3M10000019D02	Enterococcus faecalis
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167	E3M10000020G04	Enterococcus faecalis
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170	E3M10000021A11	Enterococcus faecalis
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172	E3M10000021C03	Enterococcus faecalis
173	E3M10000021C04	Enterococcus faecalis
174	E3M10000021C08	Enterococcus faecalis
175	E3M10000021D04	Enterococcus faecalis
176	E3M10000021E10	Enterococcus faecalis
177	E3M10000021G04	Enterococcus faecalis
178	E3M10000021G10	Enterococcus faecalis
179	E3M10000021G11	Enterococcus faecalis
180	E3M10000021H11	Enterococcus faecalis
181	E3M10000022A04	Enterococcus faecalis
182	E3M10000022A11	Enterococcus faecalis
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184	E3M10000022B05	Enterococcus faecalis
185	E3M10000022B07	Enterococcus faecalis
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187	E3M10000022C06	Enterococcus faecalis
188 189	E3M10000022C09	Enterococcus faecalis
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196	E3M10000023A06 E3M10000023A07	Enterococcus faecalis Enterococcus faecalis
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198	E3M10000023A09 E3M10000023B02	Enterococcus faecalis
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200	E3M10000023B06 E3M10000023C03	Enterococcus faecalis
	E3M10000023C03 E3M10000023C04	Enterococcus faecalis
202	E3M10000023C04	Enterococcus faecalis

SeqID	Clone name	Organism
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204	E3M10000023C08	Enterococcus faecalis
205	E3M10000023C09	Enterococcus faecalis
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207	E3M10000023D04	Enterococcus faecalis
208	E3M10000023D10	Enterococcus faecalis
209	E3M10000023E04	Enterococcus faecalis
210	E3M10000023E07	Enterococcus faecalis
211	E3M10000023E09	Enterococcus faecalis
212	E3M10000023F02	Enterococcus faecalis
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216	E3M10000023G10	Enterococcus faecalis
217	E3M10000023H08	Enterococcus faecalis
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220	E3M10000024A08	Enterococcus faecalis
221	E3M10000024C06	Enterococcus faecalis
222	E3M10000025A06	Enterococcus faecalis
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227	E3M10000025C01	Enterococcus faecalis
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230	E3M10000025C07	Enterococcus faecalis
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232	E3M10000025C09	Enterococcus faecalis
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247	E3M10000025G07	Enterococcus faecalis
248	E3M10000025G09	Enterococcus faecalis
249	E3M10000027A02	Enterococcus faecalis
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251	E3M10000027A09	Enterococcus faecalis

SeqID	Clone name	Organism
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253	E3M10000027B08	Enterococcus faecalis
254	E3M10000027B09	Enterococcus faecalis
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256	E3M10000027C03	Enterococcus faecalis
257	E3M10000027C08	Enterococcus faecalis
258	E3M10000027D03	Enterococcus faecalis
259	E3M10000027D05	Enterococcus faecalis
260	E3M10000027D08	Enterococcus faecalis
261	E3M10000027D10	Enterococcus faecalis
262	E3M10000027G01	Enterococcus faecalis
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266	E3M10000028A02	Enterococcus faecalis
267	E3M10000028A03	Enterococcus faecalis
268	E3M10000028A04	Enterococcus faecalis
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270	E3M10000028A06	Enterococcus faecalis
271	E3M10000028A08	Enterococcus faecalis
272	E3M10000028B01	Enterococcus faecalis
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277	E3M10000028B06	Enterococcus faecalis
278	E3M10000028B07 E3M10000028B08	Enterococcus faecalis
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280	E3M10000028C01	Enterococcus faecalis Enterococcus faecalis
282	E3M10000028C02	Enterococcus faecalis Enterococcus faecalis
283	E3M1000028C04	Enterococcus faecalis Enterococcus faecalis
284	E3M1000028C05	Enterococcus faecalis
285	E3M1000028C07	Enterococcus faecalis
286	E3M1000028C07	Enterococcus faecalis
287	E3M1000028C08	Enterococcus faecalis
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289	E3M10000028D05	Enterococcus faecalis
290	E3M1000028D06	Enterococcus faecalis
291	E3M1000028D08	Enterococcus faecalis
292	E3M10000028E01	Enterococcus faecalis
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296	E3M10000028F03	Enterococcus faecalis
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298	E3M10000028F05	Enterococcus faecalis
299	E3M10000028F06	Enterococcus faecalis
300	E3M10000028F07	Enterococcus faecalis
		

SeqID	Clone name	Organism
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307	E3M10000029A04	Enterococcus faecalis
308	E3M10000029A05	Enterococcus faecalis
309	E3M10000029A10	Enterococcus faecalis
310	E3M10000029A11	Enterococcus faecalis
311	E3M10000029B01	Enterococcus faecalis
312	E3M10000029B02	Enterococcus faecalis
313	E3M10000029B05	Enterococcus faecalis
314	E3M10000029B06	Enterococcus faecalis
315	E3M10000029B08	Enterococcus faecalis
316	E3M10000029B11	Enterococcus faecalis
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325	E3M10000029C08	Enterococcus faecalis
326	E3M10000029C09	Enterococcus faecalis
327	E3M10000029C10	Enterococcus faecalis
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329	E3M10000029D01	Enterococcus faecalis
330	E3M10000029D03	Enterococcus faecalis
331	E3M10000029D04	Enterococcus faecalis
	E3M10000029D05	Enterococcus faecalis
333	E3M10000029D06 E3M10000029D08	Enterococcus faecalis
335	E3M10000029D08	Enterococcus faecalis
336	E3M10000029E01	Enterococcus faecalis Enterococcus faecalis
337	E3M10000029E01	
338	E3M10000029E02	Enterococcus faecalis Enterococcus faecalis
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340	E3M10000029E07	Enterococcus faecalis Enterococcus faecalis
341	E3M10000029E07	Enterococcus faecalis Enterococcus faecalis
342	E3M10000029E09	Enterococcus faecalis Enterococcus faecalis
343	E3M10000029E09	Enterococcus faecalis
344	E3M10000029F01	Enterococcus faecalis Enterococcus faecalis
345	E3M10000029F05	Enterococcus faecalis Enterococcus faecalis
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345 346 347 348 349	E3M1000029F05 E3M1000029F06 E3M1000029F09 E3M1000029F10 E3M1000029F11	Enterococcus faecalis Enterococcus faecalis Enterococcus faecalis Enterococcus faecalis Enterococcus faecalis

SeqID	Clone name	Organism
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351	E3M10000029G01	Enterococcus faecalis
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353	E3M10000029G05	Enterococcus faecalis
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356	E3M10000029G09	Enterococcus faecalis
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359	E3M10000029G12	Enterococcus faecalis
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361	E3M10000029H04	Enterococcus faecalis
362	E3M10000029H05	Enterococcus faecalis
363	E3M10000029H07	Enterococcus faecalis
364	E3M10000029H08	Enterococcus faecalis
365	E3M10000029H11	Enterococcus faecalis
366	E3M10000030A05	Enterococcus faecalis
367	E3M10000030A08	Enterococcus faecalis
368	E3M10000030A09	Enterococcus faecalis
369	E3M10000030A11	Enterococcus faecalis
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372	E3M10000030B05	Enterococcus faecalis
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385	E3M10000030D09	Enterococcus faecalis
386	E3M10000030D10	Enterococcus faecalis
387	E3M10000030D12	Enterococcus faecalis
388	E3M10000030E01	Enterococcus faecalis
389	E3M10000030E02	Enterococcus faecalis
390	E3M10000030E04	Enterococcus faecalis
391	E3M10000030E08	Enterococcus faecalis
392	E3M10000030E09	Enterococcus faecalis
393	E3M10000030E10	Enterococcus faecalis
394	E3M10000030F01	Enterococcus faecalis
395	E3M10000030F04	Enterococcus faecalis
396	E3M10000030F06	Enterococcus faecalis
397	E3M10000030F07	Enterococcus faecalis
398	E3M10000030F10	Enterococcus faecalis

TABLE IA

SeqID	Clone name	Organism
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401	E3M10000030G03	Enterococcus faecalis
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403	E3M10000030G08	Enterococcus faecalis
404	E3M10000030G09	Enterococcus faecalis
405	E3M10000030G12	Enterococcus faecalis
406	E3M10000030H03	Enterococcus faecalis
407	E3M10000030H04	Enterococcus faecalis
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411	E3M10000030H10	Enterococcus faecalis
412	E3M10000030H11	Enterococcus faecalis
413	E3M10000031A02	Enterococcus faecalis
414	E3M10000031A06	Enterococcus faecalis
415	E3M10000031A07	Enterococcus faecalis
416	E3M10000031A08	Enterococcus faecalis
417	E3M10000031B02	Enterococcus faecalis
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426	E3M10000031C06	Enterococcus faecalis
427	E3M10000031C10	Enterococcus faecalis
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436	E3M10000031F04	Enterococcus faecalis
437	E3M10000031F07	Enterococcus faecalis
438	E3M10000031F09	Enterococcus faecalis
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440	E3M10000031G03	Enterococcus faecalis
441	E3M10000031G04	Enterococcus faecalis
442	E3M10000031G05	Enterococcus faecalis
443	E3M10000031G06	Enterococcus faecalis
444	E3M10000031G07	Enterococcus faecalis
445	E3M10000031G08	Enterococcus faecalis
446	E3M10000031G11	Enterococcus faecalis
447	E3M10000031H05	Enterococcus faecalis

SeqID	Clone name	Organism
448	E3M10000031H06	Enterococcus faecalis
449	E3M10000031H07	Enterococcus faecalis
450	E3M10000031H08	Enterococcus faecalis
451	E3M10000031H10	Enterococcus faecalis
452	E3M10000031H11	Enterococcus faecalis
453	E3M10000032A02	Enterococcus faecalis
454	E3M10000032A04	Enterococcus faecalis .
455	E3M10000032A06	Enterococcus faecalis
456	E3M10000032A07	Enterococcus faecalis
457	E3M10000032A08	Enterococcus faecalis
458	E3M10000032A09	Enterococcus faecalis
459	E3M10000032A10	Enterococcus faecalis
460	E3M10000032A11	Enterococcus faecalis
461	E3M10000032B03	Enterococcus faecalis
462	E3M10000032B04	Enterococcus faecalis
463	E3M10000032B07	Enterococcus faecalis
464	E3M10000032B08	Enterococcus faecalis
465	E3M10000032B09	Enterococcus faecalis
466	E3M10000032B11	Enterococcus faecalis
467	E3M10000032B12	Enterococcus faecalis
468	E3M10000032C01	Enterococcus faecalis
469	E3M10000032C02	Enterococcus faecalis
470	E3M10000032C03	Enterococcus faecalis
471	E3M10000032C04	Enterococcus faecalis
472	E3M10000032C06	Enterococcus faecalis
473	E3M10000032C09	Enterococcus faecalis
474	E3M10000032C11	Enterococcus faecalis
475	E3M10000032C12	Enterococcus faecalis
476	E3M10000032D01 E3M10000032D02	Enterococcus faecalis
477	E3M10000032D02	Enterococcus faecalis
478	E3M10000032D03	Enterococcus faecalis Enterococcus faecalis
480	E3M10000032D08	Enterococcus faecalis
481	E3M10000032D09	Enterococcus faecalis
482	E3M10000032D12	Enterococcus faecalis
483	E3M10000032E04	Enterococcus faecalis
484	E3M10000032E08	Enterococcus faecalis
485	E3M1000032E10	Enterococcus faecalis
486	E3M10000032E10	Enterococcus faecalis
487	E3M10000032E11	Enterococcus faecalis
488	E3M10000032F02	Enterococcus faecalis
489	E3M10000032F03	Enterococcus faecalis
490	E3M10000032F05	Enterococcus faecalis
491	E3M10000032F07	Enterococcus faecalis
492	E3M10000032F08	Enterococcus faecalis
493	E3M10000032F11	Enterococcus faecalis
494	E3M10000032F12	Enterococcus faecalis
495	E3M10000032G01	Enterococcus faecalis
496	E3M10000032G02	Enterococcus faecalis

SeqID	Clone name	Organism
497	E3M10000032G04	Enterococcus faecalis
498	E3M10000032G05	Enterococcus faecalis
499	E3M10000032G06	Enterococcus faecalis
500	E3M10000032G07	Enterococcus faecalis
501	E3M10000032H05	Enterococcus faecalis
502	E3M10000032H06	Enterococcus faecalis
503	E3M10000032H08	Enterococcus faecalis
504	E3M10000032H09	Enterococcus faecalis
505	E3M10000032H10	Enterococcus faecalis
506	E3M10000033A03	Enterococcus faecalis
507	E3M10000033A04	Enterococcus faecalis
508	E3M10000033A05	Enterococcus faecalis
509	E3M10000033A06	Enterococcus faecalis
510	E3M10000033A07	Enterococcus faecalis
511	E3M10000033A08	Enterococcus faecalis
512	E3M10000033A11	Enterococcus faecalis
513	E3M10000033B01	Enterococcus faecalis
514	E3M10000033B02	Enterococcus faecalis
515	E3M10000033B04	Enterococcus faecalis
516	E3M10000033B05	Enterococcus faecalis
517	E3M10000033B06	Enterococcus faecalis
518	E3M10000033B08	Enterococcus faecalis
519	E3M10000033B09	Enterococcus faecalis
520	E3M10000033C01	Enterococcus faecalis
521	E3M10000033C02	Enterococcus faecalis
522	E3M10000033C05	Enterococcus faecalis
523	E3M10000033C09	Enterococcus faecalis
524 525	E3M10000033C10	Enterococcus faecalis
526	E3M10000033C11 E3M10000033C12	Enterococcus faecalis Enterococcus faecalis
527	E3M10000033C12	Enterococcus faecalis Enterococcus faecalis
528	E3M10000033D01	Enterococcus faecalis
529	E3M10000033D04	Enterococcus faecalis
530	E3M1000033D03	Enterococcus faecalis
531	E3M1000033D00	Enterococcus faecalis
532	E3M1000033D10	Enterococcus faecalis
533	E3M10000033D11	Enterococcus faecalis
534	E3M10000033E02	Enterococcus faecalis
535	E3M10000033E03	Enterococcus faecalis
536	E3M10000033E04	Enterococcus faecalis
537	E3M10000033E05	Enterococcus faecalis
538	E3M10000033E07	Enterococcus faecalis
539	E3M10000033E08	Enterococcus faecalis
540	E3M10000033E09	Enterococcus faecalis
541	E3M10000033E11	Enterococcus faecalis
542	E3M10000033F01	Enterococcus faecalis
543	E3M10000033F03	Enterococcus faecalis
544	E3M10000033F04	Enterococcus faecalis
545	E3M10000033F05	Enterococcus faecalis
		L

SeqID	Clone name	Organism
546	E3M10000033F07	Enterococcus faecalis
547	E3M10000033F08	Enterococcus faecalis
548	E3M10000033F10	Enterococcus faecalis
549	E3M10000033F12	Enterococcus faecalis
550	E3M10000033G01	Enterococcus faecalis
551	E3M10000033G02	Enterococcus faecalis
552	E3M10000033G03	Enterococcus faecalis
553	E3M10000033G04	Enterococcus faecalis
554	E3M10000033G06	Enterococcus faecalis
555	E3M10000033G07	Enterococcus faecalis
556	E3M10000033G08	Enterococcus faecalis
557	E3M10000033G09	Enterococcus faecalis
558	E3M10000033G12	Enterococcus faecalis
559	E3M10000033H02	Enterococcus faecalis
560	E3M10000033H04	Enterococcus faecalis
561	E3M10000033H05	Enterococcus faecalis
562	E3M10000033H07	Enterococcus faecalis
563	E3M10000033H08	Enterococcus faecalis
564	E3M10000033H09	Enterococcus faecalis
565	E3M10000033H10	Enterococcus faecalis
566	E3M10000033H11	Enterococcus faecalis
567	E3M10000034A02	Enterococcus faecalis
568	E3M10000034A03	Enterococcus faecalis
569	E3M10000034A04	Enterococcus faecalis
570	E3M10000034B02	Enterococcus faecalis
571	E3M10000034B04	Enterococcus faecalis
572	E3M10000034C04	Enterococcus faecalis
573	E3M10000034D01	Enterococcus faecalis
574	E3M10000034D02	Enterococcus faecalis
575	E3M10000034E01	Enterococcus faecalis
576	E3M10000034E04	Enterococcus faecalis
577	E3M10000034F02	Enterococcus faecalis
578	E3M10000034F03	Enterococcus faecalis
579 580	E3M10000034F04 E3M10000034G02	Enterococcus faecalis
581	E3M10000034G02	Enterococcus faecalis
582	E3M10000034G03	Enterococcus faecalis Enterococcus faecalis
583	E3M10000034H02	Enterococcus faecalis Enterococcus faecalis
584	E3M10000034H03	J
585	E3M10000035A02	Enterococcus faecalis
586	E3M10000035A04	Enterococcus faecalis Enterococcus faecalis
587	E3M10000035A05	Enterococcus faecalis Enterococcus faecalis
588	E3M10000035A08	Enterococcus faecalis Enterococcus faecalis
589	E3M10000035A08	Enterococcus faecalis Enterococcus faecalis
590	E3M10000035A09	Enterococcus faecalis
591	E3M10000035A11	Enterococcus faecalis Enterococcus faecalis
592	E3M10000035B01	Enterococcus faecalis
593	E3M10000035B03	Enterococcus faecalis Enterococcus faecalis
594	E3M10000035B06	Enterococcus faecalis Enterococcus faecalis
394	/ NGCCOOODIMENT	Emerococcus jaecans

SegID Clone name Organism 595 E3M10000035B08 Enterococcus faecalis 596 E3M10000035B10 Enterococcus faecalis 597 E3M10000035B11 Enterococcus faecalis 598 E3M10000035B12 Enterococcus faecalis 500 E3M10000035C01 Enterococcus faecalis 600 E3M10000035C03 Enterococcus faecalis 601 E3M10000035C04 Enterococcus faecalis 602 E3M10000035C05 Enterococcus faecalis 603 E3M10000035C06 Enterococcus faecalis 604 E3M10000035C07 Enterococcus faecalis 605 E3M10000035C08 Enterococcus faecalis 606 E3M10000035C09 Enterococcus faecalis 607 E3M10000035C11 Enterococcus faecalis 608 E3M10000035C12 Enterococcus faecalis 609 E3M10000035D02 Enterococcus faecalis 610 E3M10000035D03 Enterococcus faecalis 611 E3M10000035D04 Enterococcus faecalis 612 E3M10000035D05 Enterococcus faecalis E3M10000035D10 613 Enterococcus faecalis 614 E3M10000035D11 Enterococcus faecalis E3M10000035E03 Enterococcus faecalis 615 616 E3M10000035E04 Enterococcus faecalis 617 E3M10000035E05 Enterococcus faecalis 618 E3M10000035E07 Enterococcus faecalis 619 E3M10000035E08 Enterococcus faecalis 620 E3M10000035E09 Enterococcus faecalis Enterococcus faecalis 621 E3M10000035E10 622 E3M10000035E11 Enterococcus faecalis 623 E3M10000035E12 Enterococcus faecalis 624 E3M10000035F01 Enterococcus faecalis 625 E3M10000035F02 Enterococcus faecalis 626 E3M10000035F03 Enterococcus faecalis 627 E3M10000035F06 Enterococcus faecalis 628 E3M10000035F07 Enterococcus faecalis 629 E3M10000035F08 Enterococcus faecalis 630 E3M10000035F09 Enterococcus faecalis 631 E3M10000035F11 Enterococcus faecalis 632 E3M10000035F12 Enterococcus faecalis 633 E3M10000035G02 Enterococcus faecalis 634 E3M10000035G04 Enterococcus faecalis 635 E3M10000035G05 Enterococcus faecalis 636 E3M10000035G08 Enterococcus faecalis 637 E3M10000035G09 Enterococcus faecalis 638 E3M10000035G10 Enterococcus faecalis 639 E3M10000035G11 Enterococcus faecalis 640 E3M10000035H03 Enterococcus faecalis 641 E3M10000035H06 Enterococcus faecalis 642 E3M10000035H09 Enterococcus faecalis

PCT/US01/09180

Enterococcus faecalis

643

E3M10000035H11

SeqID	Clone name	Organism
644	E3M10000036A03	Enterococcus faecalis
645	E3M10000036A04	Enterococcus faecalis
646	E3M10000036A05	Enterococcus faecalis
647	E3M10000036A06	Enterococcus faecalis
648	E3M10000036A07	Enterococcus faecalis
649	E3M10000036A08	Enterococcus faecalis
650	E3M10000036A09	Enterococcus faecalis
651	E3M10000036A10	Enterococcus faecalis
652	E3M10000036B01	Enterococcus faecalis
653	E3M10000036B03	Enterococcus faecalis
654	E3M10000036B06	Enterococcus faecalis
655	E3M10000036B07	Enterococcus faecalis
656	E3M10000036B08	Enterococcus faecalis
657	E3M10000036B09	Enterococcus faecalis
658	E3M10000036B11	Enterococcus faecalis
659	E3M10000036B12	Enterococcus faecalis
660	E3M10000036C01	Enterococcus faecalis
661	E3M10000036C03	Enterococcus faecalis
662	E3M10000036C06	Enterococcus faecalis
663	E3M10000036C07	Enterococcus faecalis
664	E3M10000036C08	Enterococcus faecalis
665	E3M10000036C09	Enterococcus faecalis
666	E3M10000036C10	Enterococcus faecalis
667	E3M10000036C11	Enterococcus faecalis
668	E3M10000036D03	Enterococcus faecalis
669	E3M10000036D04	Enterococcus faecalis
670	E3M10000036D06	Enterococcus faecalis
671	E3M10000036D08	Enterococcus faecalis
672	E3M10000036D09	Enterococcus faecalis
673	E3M10000036D10	Enterococcus faecalis
674	E3M10000036D11	Enterococcus faecalis
675	E3M10000036D12	Enterococcus faecalis
676	E3M10000036E01	Enterococcus faecalis
677	E3M10000036E04	Enterococcus faecalis
678	E3M10000036E05	Enterococcus faecalis
679	E3M10000036E07	Enterococcus faecalis
680	E3M10000036E08	Enterococcus faecalis
681	E3M10000036F03	Enterococcus faecalis
682	E3M10000036F04	Enterococcus faecalis
683	E3M10000036F05	Enterococcus faecalis
684	E3M10000036F08	Enterococcus faecalis
685	E3M10000036F09	Enterococcus faecalis
686	E3M10000036F10	Enterococcus faecalis
687	E3M10000036F12	Enterococcus faecalis
688	E3M1000036G01	Enterococcus faecalis
689	E3M10000036G02	Enterococcus faecalis
690	E3M10000036G03	Enterococcus faecalis
691	E3M10000036G04	Enterococcus faecalis
692	E3M10000036G06	Enterococcus faecalis

SeqID	Clone паme	Organism
693	E3M10000036G10	Enterococcus faecalis
694	ЕЗМ10000036Н02	Enterococcus faecalis
695	E3M10000036H03	Enterococcus faecalis
696	E3M10000036H04	Enterococcus faecalis
697	E3M10000036H05	Enterococcus faecalis
698	E3M10000036H06	Enterococcus faecalis
699	E3M10000036H07	Enterococcus faecalis
700	E3M10000036H08	Enterococcus faecalis
701	E3M10000036H09	Enterococcus faecalis
702	E3M10000036H10	Enterococcus faecalis
703	E3M10000037A03	Enterococcus faecalis
704	E3M10000037A06	Enterococcus faecalis
705	E3M10000037A08	Enterococcus faecalis
706	E3M10000037A09	Enterococcus faecalis
707	E3M10000037A10	Enterococcus faecalis
708	E3M10000037B02	Enterococcus faecalis
709	E3M10000037B07	Enterococcus faecalis
710	E3M10000037B08	Enterococcus faecalis
711	E3M10000037B11	Enterococcus faecalis
712	E3M10000037C01	Enterococcus faecalis
713	E3M10000037C02	Enterococcus faecalis
714	E3M10000037C04	Enterococcus faecalis
715	E3M10000037C05	Enterococcus faecalis
716	E3M10000037C07	Enterococcus faecalis
717	E3M10000037C11	Enterococcus faecalis
718	E3M10000037C12	Enterococcus faecalis
719	E3M10000037D02	Enterococcus faecalis
720	E3M10000037D03	Enterococcus faecalis
721	E3M10000037D04	Enterococcus faecalis
722	E3M10000037D05	Enterococcus faecalis
723	E3M10000037D06	Enterococcus faecalis
724	E3M10000037D09	Enterococcus faecalis
725	E3M10000037D11	Enterococcus faecalis
726 727	E3M10000037E01 E3M10000037E02	Enterococcus faecalis
727	E3M10000037E02 E3M10000037E03	Enterococcus faecalis
728	E3M10000037E03	Enterococcus faecalis
729	E3M10000037E05 E3M10000037E07	Enterococcus faecalis
730	E3M10000037E07	Enterococcus faecalis
731	E3M10000037E08	Enterococcus faecalis
733	E3M10000037E10	Enterococcus faecalis
734	E3M10000037E12 E3M10000037F01	Enterococcus faecalis Enterococcus faecalis
735	E3M10000037F01	
736	E3M10000037F02	Enterococcus faecalis
737	E3M10000037F07	Enterococcus faecalis
738	E3M10000037F07	Enterococcus faecalis Enterococcus faecalis
739	E3M10000037F12	· · · · · · · · · · · · · · · · · · ·
740	E3M10000037G01	Enterococcus faecalis
740	E3M10000037G02	Enterococcus faecalis
/41	E3M1000037G03	Enterococcus faecalis

SeqID	Clone name	Organism
742	E3M10000037G05	Enterococcus faecalis
743	E3M10000037G06	Enterococcus faecalis
744	E3M10000037G07	Enterococcus faecalis
745	E3M10000037G08	Enterococcus faecalis
746	E3M10000037G10	Enterococcus faecalis
747	E3M10000037G11	Enterococcus faecalis
748	E3M10000037H02	Enterococcus faecalis
749	E3M10000037H05	Enterococcus faecalis
750	E3M10000037H07	Enterococcus faecalis
751	E3M10000037H10	Enterococcus faecalis
752	E3M10000037H11	Enterococcus faecalis
753	E3M10000038A02	Enterococcus faecalis
754	E3M10000038A03	Enterococcus faecalis
755	E3M10000038A05	Enterococcus faecalis
756	E3M10000038A06	Enterococcus faecalis
757	E3M10000038A07	Enterococcus faecalis
758	E3M10000038A09	Enterococcus faecalis
759	E3M10000038A10	Enterococcus faecalis
760	E3M10000038A11	Enterococcus faecalis
761	E3M10000038B02	Enterococcus faecalis
762	E3M10000038B03	Enterococcus faecalis
763	E3M10000038B04	Enterococcus faecalis
764	E3M10000038B05	Enterococcus faecalis
765	E3M10000038B07	Enterococcus faecalis
766	E3M10000038B08	Enterococcus faecalis
767	E3M10000038B09	Enterococcus faecalis
768	E3M10000038B11	Enterococcus faecalis
769	E3M10000038C02	Enterococcus faecalis
770	E3M10000038C03	Enterococcus faecalis
771	E3M10000038C05	Enterococcus faecalis
772	E3M10000038C07	Enterococcus faecalis
773	E3M10000038C10	Enterococcus faecalis
774	E3M10000038C12	Enterococcus faecalis
776	E3M10000038D01 E3M10000038D02	Enterococcus faecalis
777	E3M10000038D02	Enterococcus faecalis
778	E3M10000038D08	Enterococcus faecalis Enterococcus faecalis
779	E3M10000038D08	Enterococcus faecalis Enterococcus faecalis
780	E3M10000038D10	Enterococcus faecalis Enterococcus faecalis
781	E3M10000038D11	Enterococcus faecalis Enterococcus faecalis
782	E3M10000038D12	Enterococcus faecalis Enterococcus faecalis
783	E3M10000038E02	Enterococcus faecalis
784	E3M1000038E03	Enterococcus faecalis
785	E3M10000038E04	Enterococcus faecalis
786	E3M1000038E03	Enterococcus faecalis
787	E3M1000038E07	Enterococcus faecalis
788	E3M1000038E08	Enterococcus faecalis
789	E3M1000038E11	Enterococcus faecalis
790	E3M1000038F02	Enterococcus faecalis
,,,,,		Zino, Social Jacouris

SeqID	Clone name	Organism
791	E3M10000038F05	Enterococcus faecalis
792	E3M10000038F06	Enterococcus faecalis
793	E3M10000038F07	Enterococcus faecalis
794	E3M10000038F09	- Enterococcus faecalis
795	E3M10000038F10	Enterococcus faecalis
796	E3M10000038F11	Enterococcus faecalis
797	E3M10000038G02	Enterococcus faecalis
798	E3M10000038G03	Enterococcus faecalis
799	E3M10000038G06	Enterococcus faecalis
800	E3M10000038G07	Enterococcus faecalis
801	E3M10000038G11	Enterococcus faecalis
802	E3M10000038H02	Enterococcus faecalis
803	E3M10000038H05	Enterococcus faecalis
804	E3M10000038H06	Enterococcus faecalis
805	E3M10000038H07	Enterococcus faecalis
806	E3M10000038H08	Enterococcus faecalis
807	E3M10000038H09	Enterococcus faecalis
808	E3M10000038H10	Enterococcus faecalis
809	E3M10000039A02	Enterococcus faecalis
810	E3M10000039A06	Enterococcus faecalis
811	E3M10000039A07	Enterococcus faecalis
812	E3M10000039A08	Enterococcus faecalis
813	E3M10000039A10	Enterococcus faecalis
814	E3M10000039A11	Enterococcus faecalis
815	E3M10000039B01	Enterococcus faecalis
816	E3M10000039B03	Enterococcus faecalis
817	E3M10000039B04	Enterococcus faecalis
818	E3M10000039B06	Enterococcus faecalis
819	E3M10000039B07	Enterococcus faecalis
820	E3M10000039B08	Enterococcus faecalis
821	E3M10000039B09	Enterococcus faecalis
822 823	E3M10000039B11	Enterococcus faecalis
	E3M10000039C02	Enterococcus faecalis
824 825	E3M10000039C04 E3M10000039C05	Enterococcus faecalis
825	E3M10000039C05	Enterococcus faecalis
826 827	E3M10000039C06	Enterococcus faecalis
827	E3M10000039C07	Enterococcus faecalis Enterococcus faecalis
829	E3M10000039C08	Enterococcus faecalis Enterococcus faecalis
830	E3M10000039C09	Enterococcus faecalis Enterococcus faecalis
831	E3M10000039C10	Enterococcus faecalis
832	E3M10000039D02	Enterococcus faecalis Enterococcus faecalis
833	E3M10000039D03	Enterococcus faecalis Enterococcus faecalis
834	E3M10000039D04	Enterococcus faecalis
835	E3M10000039E01	Enterococcus faecalis
836	E3M10000039E01	Enterococcus faecalis
837	E3M10000039E02	Enterococcus faecalis
838	E3M10000039E05	Enterococcus faecalis
839	E3M10000039E03	Enterococcus faecalis Enterococcus faecalis
039	ESMITO00033E01	Emerococcus Jaecans

SeqID	Clone name	Organism
840	E3M10000039E08	Enterococcus faecalis
841	E3M10000039F01	Enterococcus faecalis
842	E3M10000039F02	Enterococcus faecalis
843	E3M10000039F03	Enterococcus faecalis
844	E3M10000039F06	Enterococcus faecalis
845	E3M10000039F07	Enterococcus faecalis
846	E3M10000039F08	Enterococcus faecalis
847	E3M10000039G01	Enterococcus faecalis
848	E3M10000039G02	Enterococcus faecalis
849	E3M10000039G05	Enterococcus faecalis
850	E3M10000039G07	Enterococcus faecalis
851	E3M10000039G09	Enterococcus faecalis
852	E3M10000039G10	Enterococcus faecalis
853	E3M10000039H02	Enterococcus faecalis
854	E3M10000039H07	Enterococcus faecalis
855	E3M10000039H08	Enterococcus faecalis
856	E3M10000039H10	Enterococcus faecalis
857	E3M10000039H11	Enterococcus faecalis
858	E3M10000040A03	Enterococcus faecalis
859	E3M10000040A05	Enterococcus faecalis
860	E3M10000040A07	Enterococcus faecalis
861	E3M10000040A09	Enterococcus faecalis
862	E3M10000040A10	Enterococcus faecalis
863	E3M10000040A11	Enterococcus faecalis
864	E3M10000040B01	Enterococcus faecalis
865	E3M10000040B02	Enterococcus faecalis
866	E3M10000040B05	Enterococcus faecalis
867	E3M10000040B06	Enterococcus faecalis
868	E3M10000040B08	Enterococcus faecalis
869	E3M10000040B09	Enterococcus faecalis
870	E3M10000040B10	Enterococcus faecalis
871	E3M10000040B11	Enterococcus faecalis
872	E3M10000040B12	Enterococcus faecalis
873	E3M10000040C02	Enterococcus faecalis
874	E3M10000040C05	Enterococcus faecalis
875	E3M10000040C06	Enterococcus faecalis
876	E3M10000040C07	Enterococcus faecalis
877	E3M10000040C08	Enterococcus faecalis
878	E3M10000040C09	Enterococcus faecalis
879	E3M10000040C10	Enterococcus faecalis
880	E3M10000040C11	Enterococcus faecalis
881	E3M10000040C12	Enterococcus faecalis
882	E3M10000040D03	Enterococcus faecalis
883	E3M10000040D04	Enterococcus faecalis
884	E3M10000040D08	Enterococcus faecalis
885	E3M10000040D12	Enterococcus faecalis
886	E3M10000040E02	Enterococcus faecalis
887	E3M10000040E10	Enterococcus faecalis
888	E3M10000040E11	Enterococcus faecalis

SeqID	Clone name	Organism
889	E3M10000040E12	Enterococcus faecalis
890	E3M10000040F01	Enterococcus faecalis
891	E3M10000040F03	Enterococcus faecalis
892	E3M10000040F08	Enterococcus faecalis
893	E3M10000040F09	Enterococcus faecalis
894	E3M10000040F10	Enterococcus faecalis
895	E3M10000040G01	Enterococcus faecalis
896	E3M10000040G02	Enterococcus faecalis
897	E3M10000040G04	Enterococcus faecalis
898	E3M10000040G05	Enterococcus faecalis
899	E3M10000040G07	Enterococcus faecalis
900	E3M10000040G08	Enterococcus faecalis
901	E3M10000040G09	Enterococcus faecalis
902	E3M10000040G11	Enterococcus faecalis
903	E3M10000040H02	Enterococcus faecalis
904	E3M10000040H03	Enterococcus faecalis
905	E3M10000040H04	Enterococcus faecalis
906	E3M10000040H05	Enterococcus faecalis
907	E3M10000040H09	Enterococcus faecalis
908	E3M10000041A03	Enterococcus faecalis
909	E3M10000041A05	Enterococcus faecalis
910	E3M10000041A08	Enterococcus faecalis
911	E3M10000041A09	Enterococcus faecalis
912	E3M10000041A10	Enterococcus faecalis
913	E3M10000041A11	Enterococcus faecalis
914	E3M10000041B02	Enterococcus faecalis
915	E3M10000041B03	Enterococcus faecalis
916	E3M10000041B05	Enterococcus faecalis
917	E3M10000041B06 E3M10000041B08	Enterococcus faecalis
918	E3M10000041B08	Enterococcus faecalis
920	E3M10000041B09	Enterococcus faecalis Enterococcus faecalis
920	E3M10000041B11	Enterococcus faecalis
921	E3M10000041B11	Enterococcus faecalis
923	E3M10000041B12	Emerococcus faecalis
923	E3M10000041C01	Enterococcus faecalis
925	E3M10000041C08	Enterococcus faecalis
926	E3M10000041C09	Enterococcus faecalis
927	E3M10000041C10	Enterococcus faecalis
928	E3M10000041C11	Enterococcus faecalis
929	E3M10000041C12	Enterococcus faecalis
930	E3M10000041D02	Enterococcus faecalis
931	E3M10000041D03	Enterococcus faecalis
932	E3M10000041D04	Enterococcus faecalis
933	E3M10000041D05	Enterococcus faecalis
934	E3M10000041D06	Enterococcus faecalis
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936	E3M10000041D09	Enterococcus faecalis
937	E3M10000041D10	Enterococcus faecalis
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SeqID	Clone name	Organism
938	E3M10000041D11	Enterococcus faecalis
939	E3M10000041D12	Enterococcus faecalis
940	E3M10000041E02	Enterococcus faecalis
941	E3M10000041E03	Enterococcus faecalis
942	E3M10000041E05	Enterococcus faecalis
943	E3M10000041E07	Enterococcus faecalis
944	E3M10000041E10	Enterococcus faecalis
945	E3M10000041E11	Enterococcus faecalis
946	E3M10000041F03	Enterococcus faecalis
947	E3M10000041F05	Enterococcus faecalis
948	E3M10000041F06	Enterococcus faecalis
949	E3M10000041F07	Enterococcus faecalis
950	E3M10000041F08	Enterococcus faecalis
951	E3M10000041F09	Enterococcus faecalis
952	E3M10000041F10	Enterococcus faecalis
953	E3M10000041F11	Enterococcus faecalis
954	E3M10000041G02	Enterococcus faecalis
955	E3M10000041G03	Enterococcus faecalis
956	E3M10000041G04	Enterococcus faecalis
957	E3M10000041G06	Enterococcus faecalis
958	E3M10000041G07	Enterococcus faecalis
959	E3M10000041G08	Enterococcus faecalis
960	E3M10000041G09	Enterococcus faecalis
961	E3M10000041G10	Enterococcus faecalis
962	E3M10000041G12	Enterococcus faecalis
963	E3M10000041H04	Enterococcus faecalis
964	E3M10000041H05	Enterococcus faecalis
965	E3M10000041H06	Enterococcus faecalis
966	E3M10000041H07	Enterococcus faecalis
967	E3M10000041H08	Enterococcus faecalis
968	E3M10000041H09	Enterococcus faecalis
969	E3M10000041H10	Enterococcus faecalis
970	E3M10000041H11	Enterococcus faecalis
971	E3M10000042A03	Enterococcus faecalis
972	E3M10000042A08	Enterococcus faecalis
973 974	E3M10000042A10 E3M10000042B01	Enterococcus faecalis Enterococcus faecalis
975 976	E3M10000042B02 E3M10000042B04	Enterococcus faecalis Enterococcus faecalis
	E3M10000042B04	
977 978	E3M10000042B08	Enterococcus faecalis Enterococcus faecalis
978	E3M10000042B09	Enterococcus jaecalis Enterococcus faecalis
979	E3M10000042B10	Enterococcus jaecans Enterococcus faecalis
980	E3M10000042B11	Enterococcus jaecalis Enterococcus faecalis
981	E3M10000042C02	Enterococcus jaecalis Enterococcus faecalis
982	E3M10000042C03	Enterococcus faecalis Enterococcus faecalis
983	E3M10000042C04	Enterococcus jaecans Enterococcus faecalis
984	E3M10000042C10	Enterococcus jaecalis Enterococcus faecalis
985	E3M10000042D01	
986	E31VL1000004ZD02	Enterococcus faecalis

SeqID	Clone name	Organism
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988	E3M10000042D06	Enterococcus faecalis
989	E3M10000042D09	Enterococcus faecalis .
990	E3M10000042D11	Enterococcus faecalis
991	E3M10000042D12	Enterococcus faecalis
992	E3M10000042E05	Enterococcus faecalis
993	E3M10000042E12	Enterococcus faecalis
994	E3M10000042F11	Enterococcus faecalis
995	E3M10000042G01	Enterococcus faecalis
996	E3M10000042G05	Enterococcus faecalis
997	E3M10000042G07	Enterococcus faecalis
998	E3M10000042G08	Enterococcus faecalis
999	E3M10000042G11	Enterococcus faecalis
1000	E3M10000042G12	Enterococcus faecalis
1001	E3M10000042H06	Enterococcus faecalis
1002	E3M10000042H08	Enterococcus faecalis
1003	E3M10000042H11	Enterococcus faecalis
1004	E3M10000043A02	Enterococcus faecalis
1005	E3M10000043A03	Enterococcus faecalis
1006	E3M10000043A05	Enterococcus faecalis
1007	E3M10000043A08	Enterococcus faecalis
1008	E3M10000043A09	Enterococcus faecalis
1009	E3M10000043A10	Enterococcus faecalis
1010	E3M10000043A11	Enterococcus faecalis
1011	E3M10000043B01	Enterococcus faecalis
1012	E3M10000043B02	Enterococcus faecalis
1013	E3M10000043B03	Enterococcus faecalis
1014	E3M10000043B06 E3M10000043B08	Enterococcus faecalis
1015	E3M10000043B08	Enterococcus faecalis
1016	1	Enterococcus faecalis
1017	E3M10000043B10 E3M10000043B11	Enterococcus faecalis Enterococcus faecalis
1019	E3M10000043B11	Enterococcus faecalis Enterococcus faecalis
1019	E3M10000043B12	Enterococcus faecalis Enterococcus faecalis
1020	E3M10000043C01	Enterococcus faecalis Enterococcus faecalis
1021	E3M1000043C08	Enterococcus faecalis Enterococcus faecalis
1022	E3M10000043C09	Enterococcus faecalis
1023	E3M10000043D02	Enterococcus faecalis
1024	E3M10000043D02	Enterococcus faecalis
1025	E3M1000043D10	Enterococcus faecalis
1027	E3M10000043D12	Enterococcus faecalis
1027	E3M10000043E03	Enterococcus faecalis
1028	E3M1000043E03	Enterococcus faecalis
1030	E3M10000043E08	Enterococcus faecalis
1030	E3M10000043E10	Enterococcus faecalis
1032	E3M10000043E11	Enterococcus faecalis
1033	E3M10000043F03	Enterococcus faecalis
1034	E3M10000043F04	Enterococcus faecalis
1035	E3M10000043F06	Enterococcus faecalis
	1	

SeqID	Clone name	Organism
1036	E3M10000043F08	Enterococcus faecalis
1037	E3M10000043F10	Enterococcus faecalis
1038	E3M10000043F12	Enterococcus faecalis
1039	E3M10000043G03	Enterococcus faecalis
1040	E3M10000043G04	Enterococcus faecalis
1041	E3M10000043G05	Enterococcus faecalis
1042	E3M10000043G07	Enterococcus faecalis
1043	E3M10000043G08	Enterococcus faecalis
1044	E3M10000043G10	Enterococcus faecalis
1045	E3M10000043G11	Enterococcus faecalis
1046	E3M10000043G12	Enterococcus faecalis
1047	E3M10000043H02	Enterococcus faecalis
1048	E3M10000043H05	Enterococcus faecalis
1049	E3M10000043H08	Enterococcus faecalis
1050	E3M10000043H09	Enterococcus faecalis
1051	E3M10000043H11	Enterococcus faecalis
1052	E3M10000044C02	Enterococcus faecalis
1053	E3M10000044E01	Enterococcus faecalis
1054	K1M10000002F02	Klebsiella pneumoniae
1055	K1M10000003C01	Klebsiella pneumoniae
1056	K1M10000004F06	Klebsiella pneumoniae
1057	K1M10000007F01	Klebsiella pneumoniae
1058	K1M10000008C02	Klebsiella pneumoniae
1059	K1M10000008C10	Klebsiella pneumoniae
1060	K1M1000008G10	Klebsiella pneumoniae
1061	K1M10000009D04	Klebsiella pneumoniae
1062 1063	K1M10000013E04 K1M10000013E06	Klebsiella pneumoniae
1063	K1M10000019E06	Klebsiella pneumoniae Klebsiella pneumoniae
1064	K1M10000019D00	Klebsiella pneumoniae
1066	K1M10000020B02	Klebsiella pneumoniae
1067	K1M10000021H06	Klebsiella pneumoniae
1067	K1M10000022C10	Klebsiella pneumoniae
1069	K1M10000023E10	Klebsiella pneumoniae
1070	K1M10000023E10	Klebsiella pneumoniae
1071	K1M1000030E07	Klebsiella pneumoniae
1072	K1M10000031B11	Klebsiella pneumoniae
1073	K1M1000032E11	Klebsiella pneumoniae
1074	K1M10000033B02	Klebsiella pneumoniae
1075	K1M10000033E01	Klebsiella pneumoniae
1076	K1M10000036G08	Klebsiella pneumoniae
1077	K1M10000037D10	Klebsiella pneumoniae
1078	K1M10000038H09	Klebsiella pneumoniae
1079	К1М10000039Н03	Klebsiella pneumoniae
1080	K1M10000043C01	Klebsiella pneumoniae
1081	K1M10000043D05	Klebsiella pneumoniae
1082	K1M10000043H10	Klebsiella pneumoniae
1083	K1M10000044D05	Klebsiella pneumoniae
1084	K1M10000044D08	Klebsiella pneumoniae
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SeqID	Clone name	Organism
1085	K1M10000044E05	Klebsiella pneumoniae
1086	K1M10000044G05	Klebsiella pneumoniae
1087	K1M10000045A07	Klebsiella pneumoniae
1088	K1M10000045D10	Klebsiella pneumoniae
1089	K1M10000003D03	Klebsiella pneumoniae
1090	K1M10000010C02	Klebsiella pneumoniae
1091	K1M10000021H10	Klebsiella pneumoniae
1092	P1M1000008C06	Pseudomonas aeruginosa
1093	P1M10000008G04	Pseudomonas aeruginosa
1094	P1M10000010C03	Pseudomonas aeruginosa
1095	P1M10000014H10	Pseudomonas aeruginosa
1096	P1M10000015C06	Pseudomonas aeruginosa
1097	P1M10000015C09	Pseudomonas aeruginosa
1098	P1M10000016C04	Pseudomonas aeruginosa
1099	P1M10000018B01	Pseudomonas aeruginosa
1100	P1M10000018C01	Pseudomonas aeruginosa
1101	P1M10000018E01	Pseudomonas aeruginosa
1102	P1M10000018G01	Pseudomonas aeruginosa
1103	P1M10000019F01	Pseudomonas aeruginosa
1104	P1M10000021G03	Pseudomonas aeruginosa
1105	P1M10000021G05	Pseudomonas aeruginosa
1106	P1M10000022D09	Pseudomonas aeruginosa
1107	P1M10000024D06	Pseudomonas aeruginosa
1108	P1M10000024E06	Pseudomonas aeruginosa
1109	P1M10000024H03	Pseudomonas aeruginosa
1110	P1M10000025A06	Pseudomonas aeruginosa
1111	P1M10000025G07	Pseudomonas aeruginosa
1112	P1M10000025H07	Pseudomonas aeruginosa
1113	P1M10000026E06	Pseudomonas aeruginosa
1114	P1M10000026F04	Pseudomonas aeruginosa
1115	P1M10000026G09	Pseudomonas aeruginosa
1116	P1M10000026H02	Pseudomonas aeruginosa
1117	P1M10000026H05	Pseudomonas aeruginosa
1118	P1M10000027A06	Pseudomonas aeruginosa
1119	P1M10000027B02	Pseudomonas aeruginosa
1120	P1M10000027G05	Pseudomonas aeruginosa
1121	P1M10000028A08	Pseudomonas aeruginosa
1122	P1M10000028B01	Pseudomonas aeruginosa
1123	P1M10000028E02	Pseudomonas aeruginosa
1124	P1M10000029A09 P1M10000029G03	Pseudomonas aeruginosa
L		Pseudomonas aeruginosa
1126	P1M10000029H05 P1M10000032F04	Pseudomonas aeruginosa
1127	P1M10000032F04	Pseudomonas aeruginosa
1128	P1M10000033A02	Pseudomonas aeruginosa
1130	P1M10000033E03	Pseudomonas aeruginosa
1130	P1M10000033E03	Pseudomonas aeruginosa
1131	P1M10000033F01	Pseudomonas aeruginosa Pseudomonas aeruginosa
1132	P1M10000033G08	
1133	L IMITOOOOO SAOO	Pseudomonas aeruginosa

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SeqID	Clone name	Organism	
1134	P1M10000037B12	Pseudomonas aeruginosa	
1135	P1M10000037G12	Pseudomonas aeruginosa	
1136	P1M10000038B08	Pseudomonas aeruginosa	
1137	P1M10000038C03	Pseudomonas aeruginosa	
1138	P1M10000038C06	Pseudomonas aeruginosa	
1139	P1M10000038F04	Pseudomonas aeruginosa	
1140	P1M10000038G02	Pseudomonas aeruginosa	
1141	P1M10000039G05	Pseudomonas aeruginosa	
1142	P1M10000039G12	Pseudomonas aeruginosa	
1143	P1M10000040C01	Pseudomonas aeruginosa	
1144	P1M10000040C04	Pseudomonas aeruginosa	
1145	P1M10000040D04	Pseudomonas aeruginosa	
1146	P1M10000040D05	Pseudomonas aeruginosa	
1147	P1M10000040E10	Pseudomonas aeruginosa	
1148	P1M10000040H03	Pseudomonas aeruginosa	
1149	P1M10000041A12	Pseudomonas aeruginosa	
1150	P1M10000041B02	Pseudomonas aeruginosa	
1151	P1M10000041E01	Pseudomonas aeruginosa	
1152	P1M10000041F01	Pseudomonas aeruginosa	
1153	P1M10000042B12	Pseudomonas aeruginosa	
1154	P1M10000042E08	Pseudomonas aeruginosa	
1155	P1M10000043A03	Pseudomonas aeruginosa	
1156	P1M10000043D06	Pseudomonas aeruginosa	
1157	P1M10000044F07	Pseudomonas aeruginosa	
1158	P1M10000046B03 P1M10000046C07	Pseudomonas aeruginosa	
1160	P1M10000046C07	Pseudomonas aeruginosa	
1161	P1M10000046C09	Pseudomonas aeruginosa	
1162	P1M10000046G11	Pseudomonas aeruginosa	
1163	P1M1000043G11	Pseudomonas aeruginosa Pseudomonas aeruginosa	
1164	P1M10000047B04	Pseudomonas aeruginosa	
1165	P1M10000047E11	Pseudomonas aeruginosa	
1166	P1M10000047F07	Pseudomonas aeruginosa	
1167	P1M10000047C10	Pseudomonas aeruginosa	
1168	P1M10000049E08	Pseudomonas aeruginosa	
1169	P1M10000049G10	Pseudomonas aeruginosa	
1170	P1M10000050G11	Pseudomonas aeruginosa	
1171	P1M10000051D11	Pseudomonas aeruginosa	
1172	P1M10000051F01	Pseudomonas aeruginosa	
1173	P1M10000052C03	Pseudomonas aeruginosa	
1174	P1M10000052C12	Pseudomonas aeruginosa	
1175	P1M10000052E04	Pseudomonas aeruginosa	
1176	P1M10000053B12	Pseudomonas aeruginosa	
1177	P1M10000053C02	Pseudomonas aeruginosa	
1178	P1M10000053E07	Pseudomonas aeruginosa	
1179	P1M10000053F08	Pseudomonas aeruginosa	
1180	P1M10000055A11	Pseudomonas aeruginosa	
1181	P1M10000055C08	Pseudomonas aeruginosa	
1182	P1M10000055E05	Pseudomonas aeruginosa	

SeqID	Clone name	Organism
1183	P1M10000056C07	Pseudomonas aeruginosa
1184	P1M1000056F05	Pseudomonas aeruginosa
1185	P1M1000056F06	Pseudomonas aeruginosa
1186	P1M10000056G01	Pseudomonas aeruginosa
1187	P1M1000058B07	Pseudomonas aeruginosa
1188	P1M10000059B04	Pseudomonas aeruginosa
1189	P1M10000059B10	Pseudomonas aeruginosa
1190	P1M10000059B11	Pseudomonas aeruginosa
1191	P1M10000059D11	Pseudomonas aeruginosa
1192	P1M10000059H08	Pseudomonas aeruginosa
1193	P1M10000059H09	Pseudomonas aeruginosa
1194	P1M10000059105	Pseudomonas aeruginosa
1195	P1M10000060H02	Pseudomonas aeruginosa
1196	P1M10000060H04	Pseudomonas aeruginosa
1197	P1M10000061B04	Pseudomonas aeruginosa
1198	P1M10000061E04	Pseudomonas aeruginosa
1199	P1M10000061F04	Pseudomonas aeruginosa
1200	P1M10000062A12	Pseudomonas aeruginosa
1200	P1M1000002A12	Pseudomonas aeruginosa
1201	P1M10000032C03	Pseudomonas aeruginosa Pseudomonas aeruginosa
1202	P1M10000062C07	Pseudomonas aeruginosa
1203	P1M10000062C07	Pseudomonas aeruginosa
1204	P1M10000062D07	Pseudomonas aeruginosa
1206	P1M10000062D08	Pseudomonas aeruginosa
1207	P1M1000002E08	Pseudomonas aeruginosa
1207	P1M10000062F06	Pseudomonas aeruginosa
1209	PIM1000002G11	Pseudomonas aeruginosa
1210	P1M10000062H01	Pseudomonas aeruginosa
1211	P1M10000062H04	Pseudomonas aeruginosa
1212	P1M10000063F02	Pseudomonas aeruginosa
1213	PIM1000063G02	Pseudomonas aeruginosa
1214	P1M10000063H02	Pseudomonas aeruginosa
1215	P1M10000064A10	Pseudomonas aeruginosa
1216	P1M10000064C02	Pseudomonas aeruginosa
1217	P1M1000064C03	Pseudomonas aeruginosa
1218	P1M10000064D03	Pseudomonas aeruginosa
1219	P1M1000064E05	Pseudomonas aeruginosa
1220	P1M1000064G12	Pseudomonas aeruginosa
1221	P1M10000064H07	Pseudomonas aeruginosa
1222	P1M10000065A04	Pseudomonas aeruginosa
1223	P1M10000065B07	Pseudomonas aeruginosa
1224	P1M1000065C03	Pseudomonas aeruginosa
1225	P1M1000065C05	Pseudomonas aeruginosa
1226	P1M10000065D06	Pseudomonas aeruginosa
1227	P1M10000065F01	Pseudomonas aeruginosa
1228	P1M1000065G06	Pseudomonas aeruginosa
1229	P1M10000065H07	Pseudomonas aeruginosa
1230	PIM1000066A10	Pseudomonas aeruginosa
1231	P1M10000066A11	Pseudomonas aeruginosa

SeqID	Clone name	Organism
1232	P1M10000066F04	Pseudomonas aeruginosa
1233	P1M10000067A05	Pseudomonas aeruginosa
1234	P1M10000067A06	Pseudomonas aeruginosa
1235	P1M10000067A08	Pseudomonas aeruginosa
1236	P1M10000067C04	Pseudomonas aeruginosa
1237	P1M10000067C06	Pseudomonas aeruginosa
1238	P1M10000067D05	Pseudomonas aeruginosa
1239	P1M10000067F05	Pseudomonas aeruginosa
1240	P1M10000067G05	Pseudomonas aeruginosa
1241	P1M10000068A09	Pseudomonas aeruginosa
1242	P1M10000068D04	Pseudomonas aeruginosa
1243	P1M10000068F04	Pseudomonas aeruginosa
1244	P1M10000068F08	Pseudomonas aeruginosa
1245	P1M10000068G01	Pseudomonas aeruginosa
1246	P1M10000068H05	Pseudomonas aeruginosa
1247	P1M10000069D09	Pseudomonas aeruginosa
1248	P1M10000069G06	Pseudomonas aeruginosa
1249	P1M10000069H02	Pseudomonas aeruginosa
1250	P1M10000070A05	Pseudomonas aeruginosa
1251	P1M10000070B10	Pseudomonas aeruginosa
1252	P1M10000070C06	Pseudomonas aeruginosa
1253	P1M10000070D08	Pseudomonas aeruginosa
1254	P1M10000070E03	Pseudomonas aeruginosa
1255	P1M10000070G06	Pseudomonas aeruginosa
1256	P1M10000070G12	Pseudomonas aeruginosa
1257	P1M10000070H06	Pseudomonas aeruginosa
1258	P1M10000071A03	Pseudomonas aeruginosa
1259	P1M10000071C01	Pseudomonas aeruginosa
1260	P1M10000071E04	Pseudomonas aeruginosa
1261	P1M10000071F01	Pseudomonas aeruginosa
1262	P1M10000073A06	Pseudomonas aeruginosa
1263	P1M10000073B10	Pseudomonas aeruginosa
1264	P1M10000073D04	Pseudomonas aeruginosa
1265	P1M10000073D09	Pseudomonas aeruginosa
1266	P1M10000073G03	Pseudomonas aeruginosa
1267	P1M10000074B01	Pseudomonas aeruginosa
1268	P1M10000074B04	Pseudomonas aeruginosa
1269	P1M10000074E04	Pseudomonas aeruginosa
1270	P1M10000074E09	Pseudomonas aeruginosa
1271	P1M10000074F10	Pseudomonas aeruginosa
1272	P1M10000074G12	Pseudomonas aeruginosa
1273	P1M10000075A04	Pseudomonas aeruginosa
1274	P1M10000075B03	Pseudomonas aeruginosa
1275	P1M10000075F02	Pseudomonas aeruginosa
1276	P1M10000075G05	Pseudomonas aeruginosa
1277	P1M10000076D05	Pseudomonas aeruginosa
1278	P1M10000076D10	Pseudomonas aeruginosa
1279	P1M10000077A08	Pseudomonas aeruginosa
1280	P1M10000077C08	Pseudomonas aeruginosa

SeqID	Clone name	Organism
1281	P1M10000077E04	Pseudomonas aeruginosa
1282	P1M10000077H05	Pseudomonas aeruginosa
1283	P1M10000079A10	Pseudomonas aeruginosa
1284	P1M10000079B10	Pseudomonas aeruginosa
1285	P1M10000079C10	Pseudomonas aeruginosa
1286	P1M10000079D01	Pseudomonas aeruginosa
1287	P1M10000079D10	Pseudomonas aeruginosa
1288	P1M10000079F06	Pseudomonas aeruginosa
1289	P1M10000080B01	Pseudomonas aeruginosa
1290	P1M10000080B06	Pseudomonas aeruginosa
1291	P1M10000080C01	Pseudomonas aeruginosa
1292	P1M10000080C06	Pseudomonas aeruginosa
1293	P1M10000080E04	Pseudomonas aeruginosa
1294	P1M10000081D12	Pseudomonas aeruginosa
1295	P1M10000081G05	Pseudomonas aeruginosa
1296	P1M10000081H05	Pseudomonas aeruginosa
1297	P1M10000082A05	Pseudomonas aeruginosa
1298	P1M10000082B04	Pseudomonas aeruginosa
1299	P1M10000082C05	Pseudomonas aeruginosa
1300	P1M10000082D05	Pseudomonas aeruginosa
1301	P1M10000082E05	Pseudomonas aeruginosa
1302	P1M10000083A11	Pseudomonas aeruginosa
1303	P1M10000083B01	Pseudomonas aeruginosa
1304	P1M10000083B12	Pseudomonas aeruginosa
1305	P1M10000083C11	Pseudomonas aeruginosa
1306	P1M10000083C12	Pseudomonas aeruginosa
1307	P1M10000084A04	Pseudomonas aeruginosa
1308	P1M10000084D03	Pseudomonas aeruginosa
1309	P1M10000084E04	Pseudomonas aeruginosa
1310	P1M10000084E11	Pseudomonas aeruginosa
1311	P1M10000084F08 P1M10000085D06	Pseudomonas aeruginosa
1312		Pseudomonas aeruginosa
1313	P1M10000086A02	Pseudomonas aeruginosa
1314	P1M10000086B01 P1M10000086D02	Pseudomonas aeruginosa
1316	P1M10000086E05	Pseudomonas aeruginosa Pseudomonas aeruginosa
1317	P1M10000086E03	
1317	P1M10000087A11	Pseudomonas aeruginosa Pseudomonas aeruginosa
1319	P1M10000087C09	Pseudomonas aeruginosa Pseudomonas aeruginosa
1320	P1M10000087E04	Pseudomonas aeruginosa Pseudomonas aeruginosa
1320	P1M10000087F09	Pseudomonas aeruginosa Pseudomonas aeruginosa
1321	P1M10000087F09	Pseudomonas aeruginosa Pseudomonas aeruginosa
1323	P1M10000088D06	Pseudomonas aeruginosa
1324	P1M10000088D00	Pseudomonas aeruginosa
1325	P1M10000089D11	Pseudomonas aeruginosa
1326	P1M10000089G08	Pseudomonas aeruginosa
1327	P1M10000089G08	Pseudomonas aeruginosa Pseudomonas aeruginosa
1328	P1M1000090F06	Pseudomonas aeruginosa
1329	P1M1000090F08	Pseudomonas aeruginosa
1349	11111000050108	i seudomonas deruginosa

SegID	Clone name	Organism
1330	P1M10000091D02	Pseudomonas aeruginosa
1331	P1M10000091E09	Pseudomonas aeruginosa
1332	P1M10000091G10	Pseudomonas aeruginosa
1333	P1M10000092B02	Pseudomonas aeruginosa
1334	P1M10000092B10	Pseudomonas aeruginosa
1335	P1M10000092D09	Pseudomonas aeruginosa
1336	P1M10000092E02	Pseudomonas aeruginosa
1337	P1M10000092F05	Pseudomonas aeruginosa
1338	P1M10000093A03	Pseudomonas aeruginosa
1339	P1M10000093B09	Pseudomonas aeruginosa
1340	P1M10000093C08	Pseudomonas aeruginosa
1341	P1M10000093E09	Pseudomonas aeruginosa
1342	P1M10000093F03	Pseudomonas aeruginosa
1343	P1M10000093H07	Pseudomonas aeruginosa
1344	P1M10000094F04	Pseudomonas aeruginosa
1345	P1M10000094H03	Pseudomonas aeruginosa
1346	P1M10000095C01	Pseudomonas aeruginosa
1347	P1M10000095C09	Pseudomonas aeruginosa
1348	P1M10000095E04	Pseudomonas aeruginosa
1349	P1M10000095G04	Pseudomonas aeruginosa
1350	P1M10000096E04	Pseudomonas aeruginosa
1351	P1M10000096E12	Pseudomonas aeruginosa
1352	ID2	Pseudomonas aeruginosa
1353	4.1	Pseudomonas aeruginosa
1354	S1M10000001A05	Staphylococcus aureus
1355	S1M10000001A08	Staphylococcus aureus
1356	S1M10000001A09	Staphylococcus aureus
1357	S1M1000001A10	Staphylococcus aureus
1358	S1M10000001C06	Staphylococcus aureus
1359	S1M10000001D01	Staphylococcus aureus
1360	S1M10000001D02	Staphylococcus aureus
1361	S1M1000001D06	Staphylococcus aureus
1362	S1M10000001D07	Staphylococcus aureus
1363	S1M1000001E02	Staphylococcus aureus
1364	S1M10000001E04	Staphylococcus aureus
1365 1366	S1M10000001E05	Staphylococcus aureus
1366	S1M10000001E09	Staphylococcus aureus
1367	S1M10000001E10 S1M10000001E11	Staphylococcus aureus
1368	S1M10000001E11	Staphylococcus aureus
1370	S1M10000001F02	Staphylococcus aureus Staphylococcus aureus
1370	S1M10000001F04	Staphylococcus aureus Staphylococcus aureus
1371	S1M1000001F08	Staphylococcus aureus Staphylococcus aureus
1372	S1M1000001F09	Staphylococcus aureus Staphylococcus aureus
1374	S1M1000001F10	Staphylococcus aureus Staphylococcus aureus
1375	S1M1000001F11 S1M1000001G01	Staphylococcus aureus Staphylococcus aureus
1376	S1M1000001G01	Staphylococcus aureus
1377	S1M1000001G07	Staphylococcus aureus
1378	S1M1000001G08	Staphylococcus aureus Staphylococcus aureus
1370	011110000001010	isiapriyiococcus uureus

SeqID	Clone name	Organism
1379	S1M10000002A02	Staphylococcus aureus
1380	S1M10000002A09	Staphylococcus aureus
1381	S1M10000002A10	Staphylococcus aureus
1382	S1M10000002A12	Staphylococcus aureus
1383	S1M10000002B01	Staphylococcus aureus
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1428 SIM1000003A06 Staphylococcus aureus 1430 SIM1000003A06 Staphylococcus aureus 1431 SIM1000003A08 Staphylococcus aureus 1432 SIM1000003A10 Staphylococcus aureus 1433 SIM1000003A11 Staphylococcus aureus 1434 SIM1000003B06 Staphylococcus aureus 1435 SIM1000003B06 Staphylococcus aureus 1436 SIM1000003B08 Staphylococcus aureus 1437 SIM1000003B09 Staphylococcus aureus 1438 SIM1000003B12 Staphylococcus aureus 1439 SIM1000003C06 Staphylococcus aureus 1440 SIM1000003C07 Staphylococcus aureus 1441 SIM1000003C12 Staphylococcus aureus 1442 SIM1000003D05 Staphylococcus aureus 1443 SIM1000003D06 Staphylococcus aureus 1444 SIM1000003D06 Staphylococcus aureus 1445 SIM1000003D06 Staphylococcus aureus 1446 SIM1000003D07 Staphylococcus aureus 1447 SIM1000003D08 Staphylococcus aureus 1448 SIM1000003D09 Staphylococcus aureus 1449 SIM1000003E07 Staphylococcus aureus 1449 SIM1000003E09 Staphylococcus aureus 1449 SIM1000003E10 Staphylococcus aureus 1450 SIM1000003F06 Staphylococcus aureus 1451 SIM1000003F06 Staphylococcus aureus 1452 SIM1000003F06 Staphylococcus aureus 1453 SIM1000003F06 Staphylococcus aureus 1454 SIM1000003F06 Staphylococcus aureus 1455 SIM1000003F07 Staphylococcus aureus 1451 SIM1000003F08 Staphylococcus aureus 1452 SIM1000003F08 Staphylococcus aureus 1453 SIM1000003F08 Staphylococcus aureus 1454 SIM1000003G03 Staphylococcus aureus 1455 SIM1000003G04 Staphylococcus aureus 1456 SIM1000003G08 Staphylococcus aureus 1457 SIM1000003G08 Staphylococcus aureus 1458 SIM1000003G08 Staphylococcus aureus 1459 SIM1000003G08 Staphylococcus aureus 1450 SIM1000004A04 Staphylococcus aureus 1450 SIM1000004A04 Staphylococcus aureus 1450 SIM1000004A06 Staphylococcus aureus 1460 SIM1000004A007 Staphylococ	SeqID	Clone name	Organism
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1440 SIM1000003C10 Staphylococcus aureus 1441 SIM1000003C12 Staphylococcus aureus 1442 SIM1000003D05 Staphylococcus aureus 1443 SIM1000003D06 Staphylococcus aureus 1444 SIM1000003D08 Staphylococcus aureus 1445 SIM1000003D10 Staphylococcus aureus 1446 SIM1000003E07 Staphylococcus aureus 1447 SIM1000003E09 Staphylococcus aureus 1448 SIM1000003E10 Staphylococcus aureus 1449 SIM1000003F11 Staphylococcus aureus 1450 SIM1000003F02 Staphylococcus aureus 1451 SIM1000003F05 Staphylococcus aureus 1452 SIM1000003F06 Staphylococcus aureus 1454 SIM1000003F07 Staphylococcus aureus 1455 SIM1000003F08 Staphylococcus aureus 1455 SIM1000003F03 Staphylococcus aureus 1457 SIM1000003G03 Staphylococcus aureus 1458 SIM1000003G08 Staphylococcus aureus 1459 SIM	1438	S1M10000003C06	Staphylococcus aureus
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1476 S1M1000004C08 Staphylococcus aureus	1476	S1M1000004C08	Staphylococcus aureus

SeqID	Clone name	Organism
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1478	S1M10000004C10	Staphylococcus aureus
1479	S1M1000004C12	Staphylococcus aureus
1480	S1M10000004D01	Staphylococcus aureus
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1500	S1M10000004F12	Staphylococcus aureus
1501	S1M10000004G01	Staphylococcus aureus
1502	S1M10000004G02	Staphylococcus aureus
1503	S1M10000004G03	Staphylococcus aureus
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1505	S1M1000004G06	Staphylococcus aureus
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1508	S1M10000004G12	Staphylococcus aureus
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1510	S1M10000005A03	Staphylococcus aureus
1511	S1M10000005A05	Staphylococcus aureus
1512	S1M10000005A06	Staphylococcus aureus
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1514	S1M10000005A08	Staphylococcus aureus
1515	S1M10000005A09	Staphylococcus aureus
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1517	S1M10000005A11	Staphylococcus aureus
1518	S1M10000005B02	Staphylococcus aureus
1519	S1M10000005B04	Staphylococcus aureus
1520	S1M10000005B07	Staphylococcus aureus
1521	S1M10000005B08	Staphylococcus aureus
1522	S1M10000005B09	Staphylococcus aureus
1523	S1M10000005B12	Staphylococcus aureus
1524	S1M10000005C01	Staphylococcus aureus
1525	S1M1000005C05	Staphylococcus aureus

SeqID	Clone name	Organism
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1527	S1M10000005C09	Staphylococcus aureus
1528	S1M10000005C11	Staphylococcus aureus
1529	S1M10000005D01	Staphylococcus aureus
1530	S1M1000005D02	Staphylococcus aureus
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1534	S1M10000005D06	Staphylococcus aureus
1535	S1M10000005D07	Staphylococcus aureus
1536	S1M10000005D08	Staphylococcus aureus
1537	S1M10000005D09	Staphylococcus aureus
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1539	S1M10000005D12	Staphylococcus aureus
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1545	S1M10000005E08	Staphylococcus aureus
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1547	S1M10000005E11	Staphylococcus aureus
1548	S1M10000005E12	Staphylococcus aureus
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1550	S1M10000005F03	Staphylococcus aureus
1551	S1M10000005F04	Staphylococcus aureus
1552	S1M10000006A03	Staphylococcus aureus
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1555	S1M10000006A07	Staphylococcus aureus
1556	S1M1000006A08	Staphylococcus aureus
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1559	S1M1000006B02	Staphylococcus aureus
1560 1561	S1M10000006B03 S1M10000006B04	Staphylococcus aureus
1562	S1M10000006B07	Staphylococcus aureus
1563	S1M1000000B07	Staphylococcus aureus Staphylococcus aureus
1564	S1M10000006B10	Staphylococcus aureus Staphylococcus aureus
1565	S1M1000006B11	Staphylococcus aureus Staphylococcus aureus
1566	S1M1000000C02	Staphylococcus aureus
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1568	S1M1000000C00	Staphylococcus aureus
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1570	S1M1000000C08	Staphylococcus aureus
1571	S1M10000006D03	Staphylococcus aureus
1572	S1M10000000D05	Staphylococcus aureus
1573	S1M10000000D03	Staphylococcus aureus
1574	S1M10000006D07	Staphylococcus aureus
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SeqID	Clone name	Organism
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1576	S1M10000006E02	Staphylococcus aureus
1577	S1M10000006E03	Staphylococcus aureus
1578	S1M10000006E04	Staphylococcus aureus
1579	S1M10000006E07	Staphylococcus aureus
1580	S1M10000006E08	Staphylococcus aureus
1581	S1M10000006F01	Staphylococcus aureus
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1583	S1M10000006F03	Staphylococcus aureus
1584	S1M10000006F04	Staphylococcus aureus
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1588	S1M10000006G05	Staphylococcus aureus
1589	S1M1000006G06	Staphylococcus aureus
1590	S1M1000006G07	Staphylococcus aureus
1591	S1M1000006G09	Staphylococcus aureus
1592	S1M10000006G10	Staphylococcus aureus
1593	S1M1000006G11	Staphylococcus aureus
1594	S1M10000007A02	Staphylococcus aureus
1595	S1M10000007A03	Staphylococcus aureus
1596	S1M10000007B02	Staphylococcus aureus
1597	S1M10000007B11	Staphylococcus aureus
1598	S1M10000007C02	Staphylococcus aureus
1599	S1M10000007C04	Staphylococcus aureus
1600	S1M10000007C05	Staphylococcus aureus
1601	S1M10000007C06	Staphylococcus aureus
1602	S1M10000007C07	Staphylococcus aureus
1603	S1M1000007C08	Staphylococcus aureus
1604	S1M10000007C09	Staphylococcus aureus
1605	S1M10000007D03	Staphylococcus aureus
1606	S1M10000007D06	Staphylococcus aureus
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1608	S1M10000007D10	Staphylococcus aureus
1609	S1M10000007D11	Staphylococcus aureus
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		Staphylococcus aureus
1612	S1M10000007E07	Staphylococcus aureus
1613 1614	S1M10000007F01 S1M10000007F02	Staphylococcus aureus
1615	S1M10000007F02	Staphylococcus aureus
		Staphylococcus aureus
1616 1617	S1M10000007F08 S1M10000007F09	Staphylococcus aureus
1617	S1M10000007F09 S1M10000007F10	Staphylococcus aureus
1619	S1M10000007F10	Staphylococcus aureus
1620	S1M10000007F11	Staphylococcus aureus
1620	S1M10000007F12 S1M10000007G02	Staphylococcus aureus Staphylococcus aureus
1621	S1M1000007G02	Staphylococcus aureus Staphylococcus aureus
1623	S1M10000007G05	
1023	P.114110000007003	Staphylococcus aureus

SeqID	Clone name	Organism
1624	S1M10000007G07	Staphylococcus aureus
1625	S1M1000007G08	Staphylococcus aureus
1626	S1M10000008A03	Staphylococcus aureus
1627	S1M1000008A04	Staphylococcus aureus
1628	S1M10000008A05	Staphylococcus aureus
1629	S1M1000008A08	Staphylococcus aureus
1630	S1M10000008A09	Staphylococcus aureus
1631	S1M10000008A12 .	Staphylococcus aureus
1632	S1M10000008B03	Staphylococcus aureus
1633	S1M10000008B04	Staphylococcus aureus
1634	S1M10000008B06	Staphylococcus aureus
1635	S1M10000008B08	Staphylococcus aureus
1636	S1M10000008B09	Staphylococcus aureus
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1639	S1M10000008C06	Staphylococcus aureus
1640	S1M10000008C07	Staphylococcus aureus
1641	S1M10000008C08	Staphylococcus aureus
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1644	S1M10000008D09	Staphylococcus aureus
1645	S1M10000008E05	Staphylococcus aureus
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1647	S1M10000008E09	Staphylococcus aureus
1648	S1M10000008E10	Staphylococcus aureus
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1655	S1M10000008F10	Staphylococcus aureus
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1657	S1M1000008G02	Staphylococcus aureus
1658	S1M1000008G03	Staphylococcus aureus
1659 1660	S1M10000008G05	Staphylococcus aureus
1661	S1M1000009A02	Staphylococcus aureus
1662	S1M10000009A04 S1M10000009A07	Staphylococcus aureus
1663	S1M10000009A07 S1M10000009A08	Staphylococcus aureus
1664	S1M1000009A08 S1M10000009A09	Staphylococcus aureus
1665	S1M10000009A09	Staphylococcus aureus
1666	S1M10000009A10	Staphylococcus aureus Staphylococcus aureus
1667	S1M10000009A11 S1M10000009B01	Staphylococcus aureus Staphylococcus aureus
1668	S1M10000009B01	Staphylococcus aureus Staphylococcus aureus
1669	S1M1000009B02 S1M1000009B03	Staphylococcus aureus Staphylococcus aureus
1670	S1M1000009B03	Staphylococcus aureus Staphylococcus aureus
1671	S1M10000009B04	Staphylococcus aureus Staphylococcus aureus
1672	S1M1000009B03	Staphylococcus aureus Staphylococcus aureus
10/2	B11411000005D00	Siapnylococcus aureus

SeqID	Clone name	Organism
1673	S1M10000009B07	Staphylococcus aureus
1674	S1M10000009B10	Staphylococcus aureus
1675	S1M10000009B11	Staphylococcus aureus
1676	S1M1000009B12	Staphylococcus aureus
1677	S1M1000009C01	Staphylococcus aureus
1678	S1M1000009C02	Staphylococcus aureus
1679	S1M1000009C05	Staphylococcus aureus
1680	S1M1000009C06	Staphylococcus aureus
1681	S1M1000009C07	Staphylococcus aureus
1682	S1M10000009C08	Staphylococcus aureus
1683	S1M10000009C09	Staphylococcus aureus
1684	S1M10000009C10	Staphylococcus aureus
1685	S1M10000009C11	Staphylococcus aureus
1686	S1M10000009D01	Staphylococcus aureus
1687	S1M10000009D02	Staphylococcus aureus
1688	S1M10000009D03	Staphylococcus aureus
1689	S1M10000009D04	Staphylococcus aureus
1690	S1M10000009D05	Staphylococcus aureus
1691	S1M10000009D07	Staphylococcus aureus
1692	S1M10000009D09	Staphylococcus aureus
1693	S1M10000009D11	Staphylococcus aureus
1694	S1M1000009E02	Staphylococcus aureus
1695	S1M10000009E06	Staphylococcus aureus
1696	S1M10000009E08	Staphylococcus aureus
1697	S1M10000009E09	Staphylococcus aureus
1698	S1M10000009E11	Staphylococcus aureus
1699	S1M10000009E12	Staphylococcus aureus
1700	S1M10000009F01	Staphylococcus aureus
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1702	S1M10000009F03 S1M10000009F05	Staphylococcus aureus
1703	S1M1000009F05	Staphylococcus aureus
1704	S1M1000009F08	Staphylococcus aureus
1705	S1M1000009F07	Staphylococcus aureus
1700	S1M1000009F09	Staphylococcus aureus Staphylococcus aureus
1707	S1M10000009F10	Staphylococcus aureus
1709	S1M1000009G02	Staphylococcus aureus
1710	S1M1000009G05	Staphylococcus aureus
1711	S1M1000009G06	Staphylococcus aureus
1712	S1M1000009G07	Staphylococcus aureus
1713	S1M1000009G09	Staphylococcus aureus
1714	S1M1000009G10	Staphylococcus aureus
1715	S1M1000009G11	Staphylococcus aureus
1716	S1M1000009H01	Staphylococcus aureus
1717	S1M1000009H02	Staphylococcus aureus
1718	S1M10000009H03	Staphylococcus aureus
1719	S1M10000009H05	Staphylococcus aureus
1720	S1M1000009H07	Staphylococcus aureus
1721	S1M10000009H09	Staphylococcus aureus
		proprieta de la companya de la compa

SeqID	Clone name	Organism
1722	S1M10000009H11	Staphylococcus aureus
1723	S1M10000011A02	Staphylococcus aureus
1724	S1M10000011A03	Staphylococcus aureus
1725	S1M10000011A04	Staphylococcus aureus
1726	S1M10000011A06	Staphylococcus aureus
1727	S1M10000011B01	Staphylococcus aureus
1728	S1M10000011B02	Staphylococcus aureus
1729	S1M10000011B03	Staphylococcus aureus
1730	S1M10000011B04 .	Staphylococcus aureus
1731	S1M10000011B05	Staphylococcus aureus
1732	S1M10000011C01	Staphylococcus aureus
1733	S1M10000011C05	Staphylococcus aureus
1734	S1M10000011C06	Staphylococcus aureus
1735	S1M10000011D01	Staphylococcus aureus
1736	S1M10000011D02	Staphylococcus aureus
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1738	S1M10000011D06	Staphylococcus aureus
1739	S1M10000011E02	Staphylococcus aureus
1740	S1M10000011E03	Staphylococcus aureus
1741	S1M10000011E04	Staphylococcus aureus
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1743	S1M10000011F03	Staphylococcus aureus
1744	S1M10000011F04	Staphylococcus aureus
1745	S1M10000011F06	Staphylococcus aureus
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1747	S1M10000011G03	Staphylococcus aureus
1748	S1M10000011G04	Staphylococcus aureus
1749 1750	S1M10000011G05	Staphylococcus aureus
1751	S1M10000011G06 S1M10000011H01	Staphylococcus aureus
1752	S1M10000011H01	Staphylococcus aureus Staphylococcus aureus
1753	S1M10000011H03	
1754	S1M10000011H04	Staphylococcus aureus
1755	S1M1000012A02	Staphylococcus aureus Staphylococcus aureus
1756	S1M1000012A08	Staphylococcus aureus Staphylococcus aureus
1757	S1M10000012A08	Staphylococcus aureus
1758	S1M10000012A09	Staphylococcus aureus
1759	S1M10000012A11	Staphylococcus aureus
1760	S1M10000012311	Staphylococcus aureus
1761	S1M10000012B05	Staphylococcus aureus
1762	S1M10000012B06	Staphylococcus aureus
1763	S1M10000012B07	Staphylococcus aureus
1764	S1M10000012B11	Staphylococcus aureus
1765	S1M10000012D11	Staphylococcus aureus
1766	S1M10000012C03	Staphylococcus aureus
1767	S1M1000012C04	Staphylococcus aureus
1768	S1M10000012C05	Staphylococcus aureus
1769	S1M10000012C06	Staphylococcus aureus
1770	S1M10000012C11	Staphylococcus aureus

SeqID	Clone name	Organism
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1772	S1M10000012C12	Staphylococcus aureus
1773	S1M10000012D04	Staphylococcus aureus
1774	S1M1000012D00	Staphylococcus aureus
1775	S1M10000012D07	Staphylococcus aureus
1776	S1M1000012D09	Staphylococcus aureus
1777	S1M1000012D09	Staphylococcus aureus
1778	S1M10000012E01	Staphylococcus aureus
1779	S1M10000012E02	Staphylococcus aureus
1780	S1M10000012E04	Staphylococcus aureus
1781	S1M10000012E07	Staphylococcus aureus
1782	S1M10000012E08	Staphylococcus aureus
1783	S1M10000012E12	Staphylococcus aureus
1784	S1M10000012F04	Staphylococcus aureus
1785	S1M10000012F07	Staphylococcus aureus
1786	S1M10000012F08	Staphylococcus aureus
1787	S1M10000012F09	Staphylococcus aureus
1788	S1M10000012F10	Staphylococcus aureus
1789	S1M10000012F11	Staphylococcus aureus
1790	S1M10000012F12	Staphylococcus aureus
1791	S1M10000012G01	Staphylococcus aureus
1792	S1M10000012G02	Staphylococcus aureus
1793	S1M10000012G03	Staphylococcus aureus
1794	S1M10000012G06	Staphylococcus aureus
1795	S1M10000012G07	Staphylococcus aureus
1796	S1M10000012G08	Staphylococcus aureus
1797	S1M10000012G10	Staphylococcus aureus
1798	S1M10000012H05	Staphylococcus aureus
1799	S1M10000012H08	Staphylococcus aureus
1800	S1M10000012H09	Staphylococcus aureus
1801	S1M10000012H10	Staphylococcus aureus
1802	S1M10000012H11	Staphylococcus aureus
1803	S1M10000013A02	Staphylococcus aureus
1804	S1M10000013A03	Staphylococcus aureus
1805	S1M10000013A05	Staphylococcus aureus
1806	S1M10000013A07	Staphylococcus aureus
1807	S1M10000013A08	Staphylococcus aureus
1808	S1M10000013A09	Staphylococcus aureus
1809	S1M10000013A10	Staphylococcus aureus
1810	S1M10000013A11	Staphylococcus aureus
1811	S1M10000013A12	Staphylococcus aureus
1812	S1M10000013B02	Staphylococcus aureus
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1814	S1M10000013B04	Staphylococcus aureus
1815	S1M10000013B05	Staphylococcus aureus
1816	S1M10000013B06	Staphylococcus aureus
1817	S1M10000013B07	Staphylococcus aureus
1818	S1M10000013B09	Staphylococcus aureus
1819	S1M10000013B11	Staphylococcus aureus

SeqID	Clone name	Organism
1820	S1M10000013C03	Staphylococcus aureus
1821	S1M10000013C05	Staphylococcus aureus
1822	SIM10000013C07	Staphylococcus aureus
1823	S1M10000013C08	Staphylococcus aureus
1824	S1M10000013C09	Staphylococcus aureus
1825	S1M10000013C10	Staphylococcus aureus
1826	S1M10000013C11	Staphylococcus aureus
1827	S1M10000013C12	Staphylococcus aureus
1828	S1M10000013D08	Staphylococcus aureus
1829	S1M10000013D09	Staphylococcus aureus
1830	S1M10000013D11	Staphylococcus aureus
1831	S1M10000013E01	Staphylococcus aureus
1832	S1M10000013E02	Staphylococcus aureus
1833	S1M10000013E04	Staphylococcus aureus
1834	S1M10000013E06	Staphylococcus aureus
1835	S1M10000013E08	Staphylococcus aureus
1836	S1M10000013E09	Staphylococcus aureus
1837	S1M10000013E10	Staphylococcus aureus
1838	S1M10000013F02	Staphylococcus aureus
1839	S1M10000013F03	Staphylococcus aureus
1840	S1M10000013F06	Staphylococcus aureus
1841	S1M10000013F07	Staphylococcus aureus
1842	S1M10000013F08	Staphylococcus aureus
1843	S1M10000013F09	Staphylococcus aureus
1844	S1M10000013F12	Staphylococcus aureus
1845	S1M10000013G01	Staphylococcus aureus
1846	S1M10000013G04	Staphylococcus aureus
1847	S1M10000013G05	Staphylococcus aureus
1848	S1M10000013G06	Staphylococcus aureus
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1850	S1M10000013G10	Staphylococcus aureus
1851	S1M10000013G11	Staphylococcus aureus
1852	S1M10000013G12	Staphylococcus aureus
1853	S1M10000013H03	Staphylococcus aureus
1854	S1M10000013H04	Staphylococcus aureus
1855	S1M10000013H05	Staphylococcus aureus
1856	S1M10000013H07	Staphylococcus aureus
1857	S1M10000013H09	Staphylococcus aureus
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1859	S1M10000013H11	Staphylococcus aureus
1860	S1M10000014A02	Staphylococcus aureus
1861	S1M10000014A03	Staphylococcus aureus
1862	S1M10000014A05	Staphylococcus aureus
1863	S1M10000014A07	Staphylococcus aureus
1864	S1M10000014A08	Staphylococcus aureus
1865	S1M10000014A11	Staphylococcus aureus
1866	S1M10000014A12	Staphylococcus aureus
1867 1868	S1M10000014B01	Staphylococcus aureus
1808	S1M10000014B02	Staphylococcus aureus

SeqID	Clone name	Organism
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1870	S1M10000014B04	Staphylococcus aureus
1871	S1M10000014B05	Staphylococcus aureus
1872	S1M10000014B06	Staphylococcus aureus
1873	S1M10000014B07	Staphylococcus aureus
1874	S1M10000014B08	Staphylococcus aureus
1875	S1M10000014B10	Staphylococcus aureus
1876	S1M10000014B11	Staphylococcus aureus
1877	S1M10000014B12	Staphylococcus aureus
1878	S1M10000014C01	Staphylococcus aureus
1879	S1M10000014C05	Staphylococcus aureus
1880	S1M10000014C06	Staphylococcus aureus
1881	S1M10000014C07	Staphylococcus aureus
1882	S1M10000014C09	Staphylococcus aureus
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1885	S1M10000014C12	Staphylococcus aureus
1886	S1M10000014D03	Staphylococcus aureus
1887	S1M10000014D06	Staphylococcus aureus
1888	S1M10000014D08	Staphylococcus aureus
1889	S1M10000014D09	Staphylococcus aureus
1890	S1M10000014D10	Staphylococcus aureus
1891	S1M10000014E01	Staphylococcus aureus
1892	S1M10000014E04	Staphylococcus aureus
1893	S1M10000014E05	Staphylococcus aureus
1894	S1M10000014E07	Staphylococcus aureus
1895	S1M1000014E08	Staphylococcus aureus
1896	S1M10000014E09	Staphylococcus aureus
1897	S1M10000014E10 S1M10000014E12	Staphylococcus aureus
1898	S1M10000014E12 . S1M10000014F02	Staphylococcus aureus
1900	S1M10000014F02	Staphylococcus aureus
1900	S1M10000014F03	Staphylococcus aureus
1902	S1M10000014F05	Staphylococcus aureus Staphylococcus aureus
1902	S1M10000014F03	Staphylococcus aureus Staphylococcus aureus
1904	S1M10000014F09	Staphylococcus aureus
1904	S1M10000014F09	Staphylococcus aureus Staphylococcus aureus
1906	S1M10000014F10	Staphylococcus aureus
1907	S1M10000014G02	Staphylococcus aureus
1908	S1M1000014G04	Staphylococcus aureus
1909	S1M1000014G07	Staphylococcus aureus
1910	S1M10000014G08	Staphylococcus aureus
1911	S1M1000014G12	Staphylococcus aureus
1912	S1M10000014H02	Staphylococcus aureus
1913	S1M10000014H03	Staphylococcus aureus
1914	S1M10000014H04	Staphylococcus aureus
1915	S1M10000014H05	Staphylococcus aureus
1916	S1M10000014H06	Staphylococcus aureus
1917	S1M10000014H07	Staphylococcus aureus
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SeqID	Clone name	Organism
1918	S1M10000014H08	Staphylococcus aureus
1919	S1M10000014H11	Staphylococcus aureus
1920	S1M10000015A02	Staphylococcus aureus
1921	S1M10000015A03	Staphylococcus aureus
1922	\$1M10000015A05	Staphylococcus aureus
1923	S1M10000015A06	Staphylococcus aureus
1924	S1M10000015A09	Staphylococcus aureus
1925	S1M10000015A10	Staphylococcus aureus
1926	S1M10000015A11	Staphylococcus aureus
1927	S1M10000015A12	Staphylococcus aureus
1928	S1M10000015B02	Staphylococcus aureus
1929	S1M10000015B05	Staphylococcus aureus
1930	S1M10000015B08	Staphylococcus aureus
1931	S1M10000015B09	Staphylococcus aureus
1932	S1M10000015B10	Staphylococcus aureus
1933	S1M10000015C01	Staphylococcus aureus
1934	S1M10000015C02	Staphylococcus aureus
1935	S1M10000015C03	Staphylococcus aureus
1936	S1M10000015C05	Staphylococcus aureus
1937	S1M10000015C06	Staphylococcus aureus
1938	S1M10000015C08	Staphylococcus aureus
1939	S1M10000015C10	Staphylococcus aureus
1940	S1M10000015C12	Staphylococcus aureus
1941	S1M10000015D02	Staphylococcus aureus
1942	S1M10000015D03	Staphylococcus aureus
1943	S1M10000015D04	Staphylococcus aureus
1944	S1M10000015D05	Staphylococcus aureus
1945	S1M10000015D06	Staphylococcus aureus
1946	S1M10000015D12	Staphylococcus aureus
1947	S1M10000015E02	Staphylococcus aureus
1948	S1M10000015E03	Staphylococcus aureus
1949 1950	S1M10000015E06	Staphylococcus aureus
1950	S1M10000015E07 S1M10000015E09	Staphylococcus aureus
1951	S1M10000015E10	Staphylococcus aureus
1952	S1M10000015E10	Staphylococcus aureus
1954	S1M10000013E11	Staphylococcus aureus
1955	S1M10000013E12	Staphylococcus aureus Staphylococcus aureus
1956	S1M10000013F01	Staphylococcus aureus
1957	S1M1000015F02	Staphylococcus aureus Staphylococcus aureus
1958	S1M1000015F04	Staphylococcus aureus
1959	S1M1000015F04	Staphylococcus aureus
1960	S1M1000015F07	Staphylococcus aureus
1961	S1M1000015F07	Staphylococcus aureus
1962	S1M10000015F09	Staphylococcus aureus
1963	S1M1000015F10	Staphylococcus aureus
1964	S1M10000015F10	Staphylococcus aureus
1965	S1M10000015G02	Staphylococcus aureus
1966	S1M10000015G02	Staphylococcus aureus
1,700	0.11.10000013003	proprieto co

SeqID	Clone name	Organism
1967	S1M10000015G04	Staphylococcus aureus
1968	S1M10000015G05	Staphylococcus aureus
1969	S1M10000015G06	Staphylococcus aureus
1970	S1M10000015G07	Staphylococcus aureus
1971	S1M10000015G08	Staphylococcus aureus
1972	S1M10000015G09	Staphylococcus aureus
1973	S1M10000015G10	Staphylococcus aureus
1974	S1M10000015G11	Staphylococcus aureus
1975	S1M10000015H04	Staphylococcus aureus
1976	S1M10000015H06	Staphylococcus aureus
1977	S1M10000016A03	Staphylococcus aureus
1978	S1M10000016A04	Staphylococcus aureus
1979	S1M10000016A06	Staphylococcus aureus
1980	S1M10000016A07	Staphylococcus aureus
1981	S1M10000016A09	Staphylococcus aureus
1982	S1M10000016A10	Staphylococcus aureus
1983	S1M10000016A12	Staphylococcus aureus
1984	S1M10000016B02	Staphylococcus aureus
1985	S1M10000016B05	Staphylococcus aureus
1986	S1M10000016B06	Staphylococcus aureus
1987	S1M10000016B07	Staphylococcus aureus
1988	S1M10000016B08	Staphylococcus aureus
1989	S1M10000016B09	Staphylococcus aureus
1990	S1M10000016B10	Staphylococcus aureus
1991	S1M10000016B11	Staphylococcus aureus
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1993 1994	S1M10000016C01	Staphylococcus aureus
1994	S1M10000016C02	Staphylococcus aureus
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2007	S1M10000016D06	Staphylococcus aureus
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2012	S1M10000016E04	Staphylococcus aureus
2013	S1M10000016E05	Staphylococcus aureus
2014	S1M10000016E06	Staphylococcus aureus
2015	S1M10000016E07	Staphylococcus aureus
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SeqID	Clone name	Organism
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2018	S1M10000016E10	Staphylococcus aureus
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2020	S1M10000016E12	Staphylococcus aureus
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2024	S1M10000016F06	Staphylococcus aureus
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2027	S1M10000016F11	Staphylococcus aureus
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2035	S1M10000016H10	Staphylococcus aureus
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2037	S1M10000017A03	Staphylococcus aureus
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2039	\$1M10000017A08	Staphylococcus aureus
2040	S1M10000017A11	Staphylococcus aureus
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2053	S1M10000017C09	Staphylococcus aureus
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2062	S1M10000017E05	Staphylococcus aureus
2063	S1M10000017E08	Staphylococcus aureus
2064	S1M10000017E00	Staphylococcus aureus
2007	011-210000017.011	Dispire Cours and Cas

SeqID	Clone name	Organism
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2067	S1M10000017F05	Staphylococcus aureus
2068	S1M10000017F06	Staphylococcus aureus
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Z113	STIMITOCOCCUSETT	Staphylococcus aureus

SeqID	Clone name	Organism
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2160	S1M10000019D05	Staphylococcus aureus
2161	S1M10000019D06	Staphylococcus aureus
2162	S1M10000019D07	Staphylococcus aureus
		· · · · · · · · · · · · · · · · · · ·

2163 SIM10000019D09 Staphylococcus aureus	SeqID	Clone name	Organism
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2166 SIM10000019E02 Staphylococcus aureus 2167 SIM10000019F01 Staphylococcus aureus 2168 SIM10000019F05 Staphylococcus aureus 2169 SIM10000019F06 Staphylococcus aureus 2170 SIM10000019F08 Staphylococcus aureus 2171 SIM10000019F09 Staphylococcus aureus 2172 SIM10000019F09 Staphylococcus aureus 2173 SIM10000019F01 Staphylococcus aureus 2174 SIM1000019F01 Staphylococcus aureus 2175 SIM10000019F01 Staphylococcus aureus 2176 SIM1000019G04 Staphylococcus aureus 2177 SIM1000019G07 Staphylococcus aureus 2177 SIM1000019G07 Staphylococcus aureus 2178 SIM1000019G09 Staphylococcus aureus 2179 SIM1000019G10 Staphylococcus aureus 2179 SIM1000019G11 Staphylococcus aureus 2180 SIM1000019G11 Staphylococcus aureus 2181 SIM10000019H05 Staphylococcus aureus 2182 SIM10000019H05 Staphylococcus aureus 2183 SIM10000020A05 Staphylococcus aureus 2184 SIM10000020A06 Staphylococcus aureus 2185 SIM10000020A07 Staphylococcus aureus 2184 SIM10000020A01 Staphylococcus aureus 2185 SIM10000020A1 Staphylococcus aureus 2186 SIM10000020A1 Staphylococcus aureus 2187 SIM10000020B02 Staphylococcus aureus 2188 SIM10000020B03 Staphylococcus aureus 2198 SIM10000020B05 Staphylococcus aureus 2198 SIM10000020B05 Staphylococcus aureus 2199 SIM10000020B06 Staphylococcus aureus 2190 SIM10000020B07 Staphylococcus aureus 2191 SIM10000020B07 Staphylococcus aureus 2191 SIM10000020B09 Staphylococcus aureus 2192 SIM10000020B09 Staphylococcus aureus 2193 SIM10000020B09 Staphylococcus aureus 2194 SIM10000020B09 Staphylococcus aureus 2195 SIM10000020B09 Staphylococcus aureus 2196 SIM10000020B00 Staphylococcus aureus 2197 SIM10000020B01 Staphylococcus aureus 2198 SIM10000020B01 Staphylococcus aureus 2200 SIM10000020B00 Staphylococcus aureus 2201 SIM10000020B00 Staphylococcus aureus 2202 SIM10000020B00 Staphylococcus aureus 2203 SIM10000020B00 Staphylococcus aureus 2204 SIM10000020B00 Staphylococcus aureus 2205 SIM10000020B00 Staphylococcus aureus 2206 SIM10000020B00 Staphylococcus aureus 2207 SIM10000020B00 Staphylococcus aureus 2208 SIM10000020B01 Staphylococcus aureu	2164	S1M10000019D12	Staphylococcus aureus
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2168 SIM1000019F01 Staphylococcus aureus	2166	S1M10000019E02	Staphylococcus aureus
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2172 SIM10000019F09 Staphylococcus aureus 2173 SIM10000019F01 Staphylococcus aureus 2174 SIM10000019G04 Staphylococcus aureus 2175 SIM10000019G09 Staphylococcus aureus 2176 SIM10000019G10 Staphylococcus aureus 2177 SIM10000019G11 Staphylococcus aureus 2178 SIM10000019G11 Staphylococcus aureus 2179 SIM10000019F105 Staphylococcus aureus 2180 SIM10000019F108 Staphylococcus aureus 2181 SIM10000020A05 Staphylococcus aureus 2182 SIM10000020A06 Staphylococcus aureus 2183 SIM10000020A06 Staphylococcus aureus 2184 SIM10000020A07 Staphylococcus aureus 2185 SIM10000020A07 Staphylococcus aureus 2186 SIM10000020A07 Staphylococcus aureus 2187 SIM10000020A01 Staphylococcus aureus 2188 SIM10000020A02 Staphylococcus aureus 2189 SIM10000020B02 Staphylococcus aureus 2187 SIM10000020B03 Staphylococcus aureus 2188 SIM10000020B05 Staphylococcus aureus 2189 SIM10000020B05 Staphylococcus aureus 2190 SIM10000020B06 Staphylococcus aureus 2191 SIM10000020B07 Staphylococcus aureus 2191 SIM10000020B09 Staphylococcus aureus 2192 SIM10000020B09 Staphylococcus aureus 2193 SIM10000020B09 Staphylococcus aureus 2194 SIM10000020B09 Staphylococcus aureus 2194 SIM10000020B09 Staphylococcus aureus 2195 SIM10000020B03 Staphylococcus aureus 2196 SIM10000020B09 Staphylococcus aureus 2197 SIM10000020B09 Staphylococcus aureus 2198 SIM10000020B09 Staphylococcus aureus 2199 SIM10000020B09 Staphylococcus aureus 2191 SIM10000020B09 Staphylococcus aureus 2194 SIM10000020B09 Staphylococcus aureus 2195 SIM10000020B09 Staphylococcus aureus 2196 SIM10000020B09 Staphylococcus aureus 2197 SIM10000020B09 Staphylococcus aureus 2198 SIM10000020B09 Staphylococcus aureus 2199 SIM10000020B09 Staphylococcus aureus 2200 SIM10000020B09 Staphylococcus aureus 2201 SIM10000020B09 Staphylococcus aureus 2202 SIM10000020B09 Staphylococcus aureus 2203 SIM10000020B09 Staphylococcus aureus 2204 SIM10000020B09 Staphylococcus aureus 2205 SIM10000020B09 Staphylococcus aureus 2206 SIM10000020B09 Staphylococcus aureus 2207 SIM10000020B09 Staphylococcus aureus 2208 SIM10000020B09 Staphylo	2170	S1M10000019F06	Staphylococcus aureus
2173 SIM10000019F11 Staphylococcus aureus 2174 SIM10000019G04 Staphylococcus aureus 2175 SIM10000019G07 Staphylococcus aureus 2176 SIM10000019G09 Staphylococcus aureus 2177 SIM10000019G10 Staphylococcus aureus 2178 SIM1000019G11 Staphylococcus aureus 2179 SIM10000019H05 Staphylococcus aureus 2180 SIM10000020A05 Staphylococcus aureus 2181 SIM10000020A05 Staphylococcus aureus 2182 SIM1000020A06 Staphylococcus aureus 2183 SIM1000020A07 Staphylococcus aureus 2184 SIM1000020A11 Staphylococcus aureus 2185 SIM1000020A11 Staphylococcus aureus 2186 SIM1000020A12 Staphylococcus aureus 2187 SIM1000020B02 Staphylococcus aureus 2188 SIM1000020B03 Staphylococcus aureus 2189 SIM1000020B03 Staphylococcus aureus 2189 SIM1000020B03 Staphylococcus aureus 2190 SIM1000020B06 Staphylococcus aureus 2191 SIM1000020B07 Staphylococcus aureus 2191 SIM1000020B09 Staphylococcus aureus 2192 SIM1000020B09 Staphylococcus aureus 2193 SIM1000020B09 Staphylococcus aureus 2194 SIM1000020C09 Staphylococcus aureus 2195 SIM1000020C09 Staphylococcus aureus 2196 SIM1000020C09 Staphylococcus aureus 2197 SIM1000020D03 Staphylococcus aureus 2198 SIM1000020D04 Staphylococcus aureus 2199 SIM1000020D05 Staphylococcus aureus 2190 SIM1000020D06 Staphylococcus aureus 2191 SIM1000020D07 Staphylococcus aureus 2192 SIM1000020D09 Staphylococcus aureus 2193 SIM1000020D09 Staphylococcus aureus 2194 SIM1000020D00 Staphylococcus aureus 2195 SIM1000020D00 Staphylococcus aureus 2196 SIM1000020D00 Staphylococcus aureus 2197 SIM1000020D00 Staphylococcus aureus 2200 SIM1000020D00 Staphylococcus aureus 2201 SIM1000020D00 Staphylococcus aureus 2202 SIM1000020D00 Staphylococcus aureus 2203 SIM10000020D00 Staphylococcus aureus 2204 SIM1000020D00 Staphylococcus aureus 2205 SIM10000020D00 Staphylococcus aureus 2206 SIM10000020D00 Staphylococcus aureus 2207 SIM10000020D00 Staphylococcus aureus 2208 SIM10000020D00 Staphylococcus aureus 2209 SIM10000020D00 Staphylococcus aureus 2200 SIM10000020D00 Staphylococcus aureus 2200 SIM10000020D00 Staphylococcus aureus 2200 SIM10000020D00 S	2171	S1M10000019F08	Staphylococcus aureus
2174 SIM1000019G04 Staphylococcus aureus 2175 SIM1000019G07 Staphylococcus aureus 2176 SIM1000019G09 Staphylococcus aureus 2177 SIM10000019G10 Staphylococcus aureus 2178 SIM10000019G11 Staphylococcus aureus 2178 SIM10000019H05 Staphylococcus aureus 2180 SIM10000019H05 Staphylococcus aureus 2181 SIM10000020A05 Staphylococcus aureus 2182 SIM10000020A05 Staphylococcus aureus 2183 SIM10000020A06 Staphylococcus aureus 2184 SIM10000020A11 Staphylococcus aureus 2185 SIM10000020A11 Staphylococcus aureus 2186 SIM10000020A11 Staphylococcus aureus 2187 SIM10000020A11 Staphylococcus aureus 2188 SIM10000020B02 Staphylococcus aureus 2188 SIM10000020B03 Staphylococcus aureus 2189 SIM10000020B03 Staphylococcus aureus 2189 SIM10000020B05 Staphylococcus aureus 2190 SIM10000020B06 Staphylococcus aureus 2190 SIM10000020B07 Staphylococcus aureus 2191 SIM10000020B12 Staphylococcus aureus 2191 SIM10000020B12 Staphylococcus aureus 2193 SIM10000020B12 Staphylococcus aureus 2194 SIM10000020B12 Staphylococcus aureus 2195 SIM10000020B12 Staphylococcus aureus 2196 SIM10000020B10 Staphylococcus aureus 2197 SIM10000020B10 Staphylococcus aureus 2198 SIM10000020D0 Staphylococcus aureus 2199 SIM10000020D0 Staphylococcus aureus 2199 SIM10000020D0 Staphylococcus aureus 2199 SIM10000020D0 Staphylococcus aureus 2191 SIM10000020D0 Staphylococcus aureus 2191 SIM10000020D0 Staphylococcus aureus 2192 SIM10000020D0 Staphylococcus aureus 2193 SIM10000020D0 Staphylococcus aureus 2194 SIM10000020D0 Staphylococcus aureus 2195 SIM10000020D0 Staphylococcus aureus 2200 SIM10000020D0 Staphylococcus aureus 2201 SIM10000020D0 Staphylococcus aureus 2202 SIM10000020D0 Staphylococcus aureus 2203 SIM10000020B0 Staphylococcus aureus 2204 SIM10000020B0 Staphylococcus aureus 2205 SIM10000020B0 Staphylococcus aureus 2206 SIM10000020B0 Staphylococcus aureus 2207 SIM10000020B0 Staphylococcus aureus 2208 SIM10000020B0 Staphylococcus aureus 2209 SIM10000020B0 Staphylococcus aureus 2209 SIM10000020B0 Staphylococcus aureus 2200 SIM10000020B0 Staphylococcus aureus 2201 SIM10000		S1M10000019F09	Staphylococcus aureus
2175 SIM1000019G07 Staphylococcus aureus	2173	S1M10000019F11	Staphylococcus aureus
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2177 SIM1000019G10 Staphylococcus aureus			1 2 5
2178 SIM1000019G11 Staphylococcus aureus			<u></u>
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2181 S1M1000020A05 Staphylococcus aureus		1.	L _* · _
2182 S1M1000020A06 Staphylococcus aureus 2183 S1M1000020A07 Staphylococcus aureus 2184 S1M1000020A11 Staphylococcus aureus 2185 S1M10000020B02 Staphylococcus aureus 2186 S1M10000020B03 Staphylococcus aureus 2187 S1M10000020B03 Staphylococcus aureus 2188 S1M10000020B05 Staphylococcus aureus 2189 S1M10000020B06 Staphylococcus aureus 2190 S1M10000020B07 Staphylococcus aureus 2191 S1M10000020B09 Staphylococcus aureus 2192 S1M10000020B12 Staphylococcus aureus 2193 S1M10000020C09 Staphylococcus aureus 2194 S1M10000020C10 Staphylococcus aureus 2195 S1M10000020C11 Staphylococcus aureus 2196 S1M10000020D03 Staphylococcus aureus 2197 S1M10000020D04 Staphylococcus aureus 2198 S1M10000020D07 Staphylococcus aureus 2200 S1M10000020D08 Staphylococcus aureus 2201			
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2186 \$IM1000020B02 \$Staphylococcus aureus 2187 \$IM1000020B03 \$Staphylococcus aureus 2188 \$IM10000020B05 \$Staphylococcus aureus 2189 \$IM10000020B06 \$Staphylococcus aureus 2190 \$IM10000020B07 \$Staphylococcus aureus 2191 \$IM1000020B09 \$Staphylococcus aureus 2192 \$IM1000020B12 \$Staphylococcus aureus 2193 \$SIM10000020C09 \$Staphylococcus aureus 2194 \$SIM10000020C10 \$Staphylococcus aureus 2195 \$SIM10000020C11 \$Staphylococcus aureus 2196 \$IM10000020D03 \$Staphylococcus aureus 2197 \$SIM1000020D04 \$Staphylococcus aureus 2198 \$SIM10000020D06 \$Staphylococcus aureus 2199 \$SIM10000020D07 \$Staphylococcus aureus 2200 \$SIM10000020D08 \$Staphylococcus aureus 2201 \$SIM10000020D01 \$Staphylococcus aureus 2202 \$SIM10000020D03 \$Staphylococcus aureus 2204 \$SIM10000020E03 \$Staphylococcus aureus <			
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2243 \$\text{SIM10000021C04} \$\text{Staphylococcus aureus}\$ 2244 \$\text{SIM10000021C07} \$\text{Staphylococcus aureus}\$ 2245 \$\text{SIM10000021C08} \$\text{Staphylococcus aureus}\$ 2246 \$\text{SIM10000021C10} \$\text{Staphylococcus aureus}\$ 2247 \$\text{SIM10000021C11} \$\text{Staphylococcus aureus}\$ 2248 \$\text{SIM10000021C11} \$\text{Staphylococcus aureus}\$ 2249 \$\text{SIM10000021C12} \$\text{Staphylococcus aureus}\$ 2250 \$\text{SIM10000021D01} \$\text{Staphylococcus aureus}\$ 2251 \$\text{SIM10000021D03} \$\text{Staphylococcus aureus}\$ 2252 \$\text{SIM10000021D04} \$\text{Staphylococcus aureus}\$ 2253 \$\text{SIM10000021D06} \$\text{Staphylococcus aureus}\$ 2254 \$\text{SIM10000021D09} \$\text{Staphylococcus aureus}\$ 2255 \$\text{SIM10000021D10} \$\text{Staphylococcus aureus}\$ 2256 \$\text{SIM10000021E02} \$\text{Staphylococcus aureus}\$ 2258 \$\text{SIM10000021E03} \$\text{Staphylococcus aureus}\$ 2259 \$\text{SIM10000021E05} \$Staphylococcus aure	L		l 1 7
2244 S1M10000021C05 Staphylococcus aureus 2245 S1M10000021C07 Staphylococcus aureus 2246 S1M10000021C08 Staphylococcus aureus 2247 S1M10000021C10 Staphylococcus aureus 2248 S1M10000021C11 Staphylococcus aureus 2249 S1M10000021D01 Staphylococcus aureus 2250 S1M10000021D01 Staphylococcus aureus 2251 S1M10000021D03 Staphylococcus aureus 2252 S1M10000021D04 Staphylococcus aureus 2253 S1M10000021D06 Staphylococcus aureus 2254 S1M10000021D09 Staphylococcus aureus 2255 S1M10000021D10 Staphylococcus aureus 2256 S1M10000021E01 Staphylococcus aureus 2257 S1M10000021E02 Staphylococcus aureus 2258 S1M10000021E03 Staphylococcus aureus 2259 S1M10000021E05 Staphylococcus aureus			, - <u></u>
2245 S1M10000021C07 Staphylococcus aureus 2246 S1M10000021C08 Staphylococcus aureus 2247 S1M10000021C10 Staphylococcus aureus 2248 S1M10000021C11 Staphylococcus aureus 2249 S1M10000021C12 Staphylococcus aureus 2250 S1M10000021D01 Staphylococcus aureus 2251 S1M10000021D03 Staphylococcus aureus 2252 S1M10000021D04 Staphylococcus aureus 2253 S1M10000021D06 Staphylococcus aureus 2254 S1M10000021D09 Staphylococcus aureus 2255 S1M10000021D10 Staphylococcus aureus 2256 S1M10000021E01 Staphylococcus aureus 2257 S1M10000021E02 Staphylococcus aureus 2258 S1M10000021E03 Staphylococcus aureus 2259 S1M10000021E05 Staphylococcus aureus			
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2251 \$\text{S1M10000021D03}\$ \$\text{Staphylococcus aureus}\$ 2252 \$\text{S1M10000021D04}\$ \$\text{Staphylococcus aureus}\$ 2253 \$\text{S1M10000021D06}\$ \$\text{Staphylococcus aureus}\$ 2254 \$\text{S1M10000021D09}\$ \$\text{Staphylococcus aureus}\$ 2255 \$\text{S1M10000021D10}\$ \$\text{Staphylococcus aureus}\$ 2256 \$\text{S1M10000021E01}\$ \$\text{Staphylococcus aureus}\$ 2257 \$\text{S1M10000021E02}\$ \$\text{Staphylococcus aureus}\$ 2258 \$\text{S1M10000021E03}\$ \$\text{Staphylococcus aureus}\$ 2259 \$\text{S1M10000021E05}\$ \$\text{Staphylococcus aureus}\$			
2252 \$\text{S1M10000021D04}\$ \$\text{Staphylococcus aureus}\$ 2253 \$\text{S1M10000021D06}\$ \$\text{Staphylococcus aureus}\$ 2254 \$\text{S1M10000021D09}\$ \$\text{Staphylococcus aureus}\$ 2255 \$\text{S1M10000021D10}\$ \$\text{Staphylococcus aureus}\$ 2256 \$\text{S1M10000021E01}\$ \$\text{Staphylococcus aureus}\$ 2257 \$\text{S1M10000021E02}\$ \$\text{Staphylococcus aureus}\$ 2258 \$\text{S1M10000021E03}\$ \$\text{Staphylococcus aureus}\$ 2259 \$\text{S1M10000021E05}\$ \$\text{Staphylococcus aureus}\$	2251		
2253 \$1M1000021D06 \$Staphylococcus aureus 2254 \$1M1000021D09 \$Staphylococcus aureus 2255 \$1M1000021D10 \$Staphylococcus aureus 2256 \$1M1000021E01 \$Staphylococcus aureus 2257 \$1M1000021E02 \$Staphylococcus aureus 2258 \$1M1000021E03 \$Staphylococcus aureus 2259 \$1M1000021E05 \$Staphylococcus aureus	2252		
2254 S1M10000021D09 Staphylococcus aureus 2255 S1M10000021D10 Staphylococcus aureus 2256 S1M10000021E01 Staphylococcus aureus 2257 S1M10000021E02 Staphylococcus aureus 2258 S1M10000021E03 Staphylococcus aureus 2259 S1M10000021E05 Staphylococcus aureus	2253		
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2256 \$1M1000021E01 \$Staphylococcus aureus 2257 \$1M1000021E02 \$Staphylococcus aureus 2258 \$1M10000021E03 \$Staphylococcus aureus 2259 \$1M10000021E05 \$Staphylococcus aureus	2255	S1M10000021D10	
2258 S1M10000021E03 Staphylococcus aureus 2259 S1M10000021E05 Staphylococcus aureus	2256	S1M10000021E01	
2258 S1M10000021E03 Staphylococcus aureus 2259 S1M10000021E05 Staphylococcus aureus	2257	S1M10000021E02	Staphylococcus aureus
	2258	S1M10000021E03	Staphylococcus aureus
2260 S1M10000021E06 Staphylococcus aureus	2259	S1M10000021E05	Staphylococcus aureus
	2260	S1M10000021E06	Staphylococcus aureus

SeqID	Clone name	Organism
2261	S1M10000021E09	Staphylococcus aureus
2262	S1M10000021E12	Staphylococcus aureus
2263	S1M10000021F02	Staphylococcus aureus
2264	S1M10000021F04	Staphylococcus aureus
2265	S1M10000021F05	Staphylococcus aureus
2266	S1M10000021F06	Staphylococcus aureus
2267	S1M10000021F07	Staphylococcus aureus
2268	S1M10000021F09	Staphylococcus aureus .
2269	S1M10000021F11	Staphylococcus aureus
2270	S1M10000021G01	Staphylococcus aureus
2271	S1M10000021G03	Staphylococcus aureus
2272	S1M10000021G08	Staphylococcus aureus
2273	S1M10000021H04	Staphylococcus aureus
2274	S1M10000021H05	Staphylococcus aureus
2275	S1M10000021H07	Staphylococcus aureus
2276	S1M10000021H08	Staphylococcus aureus
2277	S1M10000021H11	Staphylococcus aureus
2278	S1M10000022A02	Staphylococcus aureus
2279	S1M10000022A03	Staphylococcus aureus
2280	S1M10000022A05	Staphylococcus aureus
2281	S1M10000022A08	Staphylococcus aureus
2282	S1M10000022A09	Staphylococcus aureus
2283	S1M10000022A12	Staphylococcus aureus
2284	S1M10000022B02	Staphylococcus aureus
2285	S1M10000022B03	Staphylococcus aureus
2286	S1M10000022B05	Staphylococcus aureus
2287	S1M10000022B06	Staphylococcus aureus
2288	S1M10000022B08	Staphylococcus aureus
2289	S1M10000022B09 S1M10000022B10	Staphylococcus aureus
2290	S1M10000022B10	Staphylococcus aureus
2291	S1M10000022B11	Staphylococcus aureus Staphylococcus aureus
2292	S1M1000022B12 S1M10000022C02	1 ^ -
2294	S1M1000022C02	Staphylococcus aureus Staphylococcus aureus
2295	S1M1000022C03	Staphylococcus aureus
2296	S1M1000022C06	Staphylococcus aureus
2297	S1M1000022C07	Staphylococcus aureus
2298	S1M1000022C07	Staphylococcus aureus
2299	S1M10000022C00	Staphylococcus aureus
2300	S1M10000022D03	Staphylococcus aureus
2301	S1M10000022D05	Staphylococcus aureus
2302	S1M10000022D06	Staphylococcus aureus
2303	S1M10000022D07	Staphylococcus aureus
2304	S1M10000022D08	Staphylococcus aureus
2305	S1M10000022D09	Staphylococcus aureus
2306	S1M10000022D11	Staphylococcus aureus
2307	S1M10000022E01	Staphylococcus aureus
2308	S1M10000022E03	Staphylococcus aureus
2309	S1M10000022E05	Staphylococcus aureus
		<u> </u>

/0955		TABLE IA	PC 1/US01/09180
SeqID	Clone name	Organism	
2310	S1M10000022E09	Staphylococcus aureus	
2311	S1M10000022F04	Staphylococcus aureus	
2312	S1M10000022F06	Staphylococcus aureus	·
2313	S1M10000022F07	Staphylococcus aureus	
2314	S1M10000022F08	Staphylococcus aureus	
2315	S1M10000022F11	Staphylococcus aureus	
2316	S1M10000022G03	Staphylococcus aureus	
2317	S1M10000022G04	Staphylococcus aureus	
2318	S1M10000022G07	Staphylococcus aureus	
2319	S1M10000022G08	Staphylococcus aureus	· · · · · · · · · · · · · · · · · · ·
2320	S1M10000022G12	Staphylococcus aureus	· · · · · · · · · · · · · · · · · · ·
2321	S1M10000022H03	Staphylococcus aureus	
2322	S1M10000022H05	Staphylococcus aureus	
2323	S1M10000022H06	Staphylococcus aureus	
2324	S1M10000022H07	Staphylococcus aureus	
2325	S1M10000022H08	Staphylococcus aureus	
2326	S1M10000022H11	Staphylococcus aureus	
2327	S1M10000023A05	Staphylococcus aureus	
2328	S1M10000023A09	Staphylococcus aureus	
2329	S1M10000023A11	Staphylococcus aureus	
2330	S1M10000023A12	Staphylococcus aureus	
2331	S1M10000023B01	Staphylococcus aureus	
2332	S1M10000023B03	Staphylococcus aureus	
2333	S1M10000023B07	Staphylococcus aureus	
2334	S1M10000023B08	Staphylococcus aureus	
2335	S1M10000023B09	Staphylococcus aureus	
2336	\$1M10000023B10	Staphylococcus aureus	
2337	S1M10000023B11	Staphylococcus aureus	
2338	S1M10000023B12	Staphylococcus aureus	
2339	S1M10000023C02	Staphylococcus aureus	
2340	S1M10000023C10	Staphylococcus aureus	
2341	S1M10000023C11	Staphylococcus aureus	
2342	S1M10000023C12	Staphylococcus aureus	
2343	S1M10000023D01	Staphylococcus aureus	
2344	S1M10000023D03	Staphylococcus aureus	
2345	S1M10000023D04	Staphylococcus aureus	
2346	S1M10000023D07	Staphylococcus aureus	
2347	S1M10000023D08	Staphylococcus aureus	
2348	S1M10000023D09	Staphylococcus aureus	
2349	S1M10000023D10	Staphylococcus aureus	
2350	S1M10000023D12	Staphylococcus aureus	
2351	S1M10000023E01	Staphylococcus aureus	
2352	S1M10000023E04	Staphylococcus aureus	
2353	S1M10000023E07	Staphylococcus aureus	
2354	S1M10000023E10	Staphylococcus aureus	
2355	S1M10000023E11	Staphylococcus aureus	
2356	S1M10000023F04	Staphylococcus aureus	
2357	S1M10000023F07	Staphylococcus aureus	
2358	S1M10000023F08	Staphylococcus aureus	

2359 S1M10000023F10 Staphylococcus aureus 2361 S1M10000023F11 Staphylococcus aureus 2362 S1M10000023G02 Staphylococcus aureus 2363 S1M10000023G03 Staphylococcus aureus 2364 S1M10000023G06 Staphylococcus aureus 2365 S1M10000023G07 Staphylococcus aureus 2365 S1M10000023G07 Staphylococcus aureus 2365 S1M10000023G08 Staphylococcus aureus 2366 S1M10000023G08 Staphylococcus aureus 2367 S1M10000023G09 Staphylococcus aureus 2368 S1M10000023G11 Staphylococcus aureus 2368 S1M10000023G11 Staphylococcus aureus 2379 S1M10000023H02 Staphylococcus aureus 2370 S1M10000023H07 Staphylococcus aureus 2371 S1M10000023H07 Staphylococcus aureus 2372 S1M10000023H07 Staphylococcus aureus 2373 S1M10000023H07 Staphylococcus aureus 2374 S1M10000023H07 Staphylococcus aureus 2375 S1M10000023H07 Staphylococcus aureus 2376 S1M10000023H07 Staphylococcus aureus 2377 S1M10000024A02 Staphylococcus aureus 2378 S1M10000024A04 Staphylococcus aureus 2379 S1M10000024A04 Staphylococcus aureus 2379 S1M10000024A04 Staphylococcus aureus 2379 S1M10000024A08 Staphylococcus aureus 2379 S1M10000024A08 Staphylococcus aureus 2379 S1M10000024A08 Staphylococcus aureus 2380 S1M10000024B08 Staphylococcus aureus 2381 S1M10000024B06 Staphylococcus aureus 2382 S1M10000024B06 Staphylococcus aureus 2383 S1M10000024B06 Staphylococcus aureus 2383 S1M10000024B06 Staphylococcus aureus 2384 S1M10000024B06 Staphylococcus aureus 2385 S1M10000024D02 Staphylococcus aureus 2386 S1M10000024C02 Staphylococcus aureus 2387 S1M10000024D03 Staphylococcus aureus 2388 S1M10000024C04 Staphylococcus aureus 2389 S1M10000024D04 Staphylococcus aureus 2380 S1M10000024D05 Staphylococcus aureus 2381 S1M10000024C07 Staphylococcus aureus 2382 S1M10000024C07 Staphylococcus aureus 2383 S1M10000024C07 Staphylococcus aureus 2390 S1M10000024C07 Staphylococcus aureus 2391 S1M10000024C07 Staphylococcus aureus 2392 S1M10000024C07 Staphylococcus aureus 2393 S1M10000024C07 Staphylococcus aureus 2394 S1M10000024C07 Staphylococcus aureus 2395 S1M10000024C07 Staphylococcus aureus 2396 S1M10000024C07 Staphyloco	SeqID	Clone name	Organism
2360 SIM10000023F11 Staphylococcus aureus		1	
2361 SIM10000023G02 Staphylococcus aureus			
2362 SIM10000023G02 Staphylococcus aureus 2363 SIM10000023G06 Staphylococcus aureus 2364 SIM10000023G06 Staphylococcus aureus 2365 SIM10000023G08 Staphylococcus aureus 2366 SIM10000023G08 Staphylococcus aureus 2367 SIM10000023G08 Staphylococcus aureus 2368 SIM10000023G01 Staphylococcus aureus 2368 SIM10000023G11 Staphylococcus aureus 2370 SIM10000023H02 Staphylococcus aureus 2371 SIM10000023H06 Staphylococcus aureus 2372 SIM10000023H07 Staphylococcus aureus 2373 SIM10000023H07 Staphylococcus aureus 2373 SIM10000023H09 Staphylococcus aureus 2374 SIM10000023H09 Staphylococcus aureus 2375 SIM10000024A02 Staphylococcus aureus 2376 SIM10000024A02 Staphylococcus aureus 2377 SIM10000024A03 Staphylococcus aureus 2378 SIM10000024A04 Staphylococcus aureus 2379 SIM10000024A08 Staphylococcus aureus 2379 SIM10000024A08 Staphylococcus aureus 2379 SIM10000024A08 Staphylococcus aureus 2378 SIM10000024A08 Staphylococcus aureus 2380 SIM10000024A08 Staphylococcus aureus 2381 SIM10000024B06 Staphylococcus aureus 2382 SIM10000024B08 Staphylococcus aureus 2383 SIM10000024B08 Staphylococcus aureus 2383 SIM10000024B08 Staphylococcus aureus 2384 SIM10000024C04 Staphylococcus aureus 2385 SIM10000024C02 Staphylococcus aureus 2386 SIM10000024C02 Staphylococcus aureus 2387 SIM10000024C03 Staphylococcus aureus 2388 SIM10000024C03 Staphylococcus aureus 2389 SIM10000024C03 Staphylococcus aureus 2389 SIM10000024C03 Staphylococcus aureus 2390 SIM10000024D03 Staphylococcus aureus 2391 SIM10000024D03 Staphylococcus aureus 2392 SIM10000024D03 Staphylococcus aureus 2393 SIM10000024E03 Staphylococcus aureus 2393 SIM10000024E03 Staphylococcus aureus 2393 SIM10000024E03 Staphylococcus aureus 2394 SIM10000024E03 Staphylococcus aureus 2395 SIM10000024E03 Staphylococcus aureus 2396 SIM10000024E03 Staphylococcus aureus 2397 SIM10000024E03 Staphylococcus aureus 2398 SIM10000024E03 Staphylococcus aureus 2399 SIM10000024E03 Staphylococcus aureus 2399 SIM10000024E03 Staphylococcus aureus 2399 SIM10000024E03 Staphylococcus aureus 2399 SIM10000024E03 Staphyloco			
2363 SIM10000023G06 Staphylococcus aureus			1 - 2 - 2
2364 S1M1000023G06 Staphylococcus aureus 2365 S1M10000023G07 Staphylococcus aureus 2366 S1M10000023G09 Staphylococcus aureus 2367 S1M10000023G09 Staphylococcus aureus 2368 S1M10000023G09 Staphylococcus aureus 2369 S1M10000023H02 Staphylococcus aureus 2369 S1M10000023H02 Staphylococcus aureus 2370 S1M10000023H07 Staphylococcus aureus 2371 S1M10000023H07 Staphylococcus aureus 2372 S1M10000023H09 Staphylococcus aureus 2373 S1M10000023H09 Staphylococcus aureus 2374 S1M10000024H09 Staphylococcus aureus 2375 S1M10000024A02 Staphylococcus aureus 2376 S1M10000024A04 Staphylococcus aureus 2377 S1M10000024A07 Staphylococcus aureus 2378 S1M10000024A08 Staphylococcus aureus 2379 S1M10000024A08 Staphylococcus aureus 2379 S1M1000024A08 Staphylococcus aureus 2380 S1M1000024B06 Staphylococcus aureus 2381 S1M1000024B06 Staphylococcus aureus 2382 S1M1000024B08 Staphylococcus aureus 2382 S1M1000024B09 Staphylococcus aureus 2383 S1M1000024B09 Staphylococcus aureus 2384 S1M1000024B09 Staphylococcus aureus 2385 S1M1000024C02 Staphylococcus aureus 2386 S1M1000024C04 Staphylococcus aureus 2387 S1M1000024C04 Staphylococcus aureus 2388 S1M1000024C04 Staphylococcus aureus 2389 S1M1000024C07 Staphylococcus aureus 2389 S1M1000024C08 Staphylococcus aureus 2391 S1M1000024E06 Staphylococcus aureus 2392 S1M1000024E06 Staphylococcus aureus 2393 S1M1000024E06 Staphylococcus aureus 2394 S1M1000024E06 Staphylococcus aureus 2395 S1M1000024E06 Staphylococcus aureus 2396 S1M1000024E07 Staphylococcus aureus 2397 S1M1000024E06 Staphylococcus aureus 2398 S1M1000024E06 Staphylococcus aureus 2399 S1M1000024E08 Staphylococcus aureus 2399 S1M1000024E08 Staphylococcus aureus 2399 S1M1000024E08 Staphylococcus aureus 2399 S1M1000024E08 Staphylococcus aureus 2400 S1M1000024G08 Staphylococcus aureus 2401			
2365 SIM10000023G07 Staphylococcus aureus			
2366 SIM10000023G08 Staphylococcus aureus			
2367 SIM10000023G99 Staphylococcus aureus			
2368 SIM10000023G11 Staphylococcus aureus			
2369 SIM10000023H02 Staphylococcus aureus			
2370 SIM10000023H06 Staphylococcus aureus			
2371 SIM10000023H09 Staphylococcus aureus			
2372 SIM10000023H09 Staphylococcus aureus		1	
Staphylococcus aureus			
2374 SIM1000024A02 Staphylococcus aureus		f	
2375 SIM10000024A04 Staphylococcus aureus			1
2376 S1M10000024A07 Staphylococcus aureus 2377 S1M10000024A08 Staphylococcus aureus 2378 S1M10000024B05 Staphylococcus aureus 2379 S1M10000024B06 Staphylococcus aureus 2380 S1M10000024B08 Staphylococcus aureus 2381 S1M10000024B08 Staphylococcus aureus 2382 S1M10000024B10 Staphylococcus aureus 2383 S1M10000024C02 Staphylococcus aureus 2384 S1M10000024C02 Staphylococcus aureus 2385 S1M10000024C07 Staphylococcus aureus 2386 S1M10000024D02 Staphylococcus aureus 2387 S1M10000024D02 Staphylococcus aureus 2388 S1M10000024D03 Staphylococcus aureus 2390 S1M10000024E01 Staphylococcus aureus 2391 S1M10000024E03 Staphylococcus aureus 2392 S1M10000024E05 Staphylococcus aureus 2393 S1M10000024E06 Staphylococcus aureus 2394 S1M10000024F03 Staphylococcus aureus 2395 <td></td> <td></td> <td></td>			
2377 SIM10000024A08 Staphylococcus aureus			
2378 S1M1000024A11 Staphylococcus aureus			1 - 2
2379 S1M10000024B05 Staphylococcus aureus 2380 S1M10000024B06 Staphylococcus aureus 2381 S1M10000024B09 Staphylococcus aureus 2382 S1M10000024B09 Staphylococcus aureus 2383 S1M10000024B00 Staphylococcus aureus 2384 S1M10000024C02 Staphylococcus aureus 2385 S1M10000024C07 Staphylococcus aureus 2386 S1M10000024D02 Staphylococcus aureus 2387 S1M10000024D03 Staphylococcus aureus 2389 S1M10000024D03 Staphylococcus aureus 2390 S1M10000024D01 Staphylococcus aureus 2391 S1M10000024E03 Staphylococcus aureus 2392 S1M10000024E03 Staphylococcus aureus 2393 S1M10000024E06 Staphylococcus aureus 2394 S1M10000024E06 Staphylococcus aureus 2395 S1M10000024E08 Staphylococcus aureus 2396 S1M10000024F03 Staphylococcus aureus 2398 S1M10000024F03 Staphylococcus aureus 2399 <td></td> <td></td> <td></td>			
2380 S1M10000024B06 Staphylococcus aureus 2381 S1M10000024B09 Staphylococcus aureus 2382 S1M10000024B10 Staphylococcus aureus 2383 S1M10000024C02 Staphylococcus aureus 2384 S1M10000024C04 Staphylococcus aureus 2385 S1M10000024C07 Staphylococcus aureus 2386 S1M10000024D02 Staphylococcus aureus 2387 S1M10000024D03 Staphylococcus aureus 2389 S1M10000024D03 Staphylococcus aureus 2390 S1M10000024D11 Staphylococcus aureus 2391 S1M10000024E03 Staphylococcus aureus 2392 S1M10000024E03 Staphylococcus aureus 2393 S1M10000024E06 Staphylococcus aureus 2394 S1M10000024E06 Staphylococcus aureus 2395 S1M10000024E08 Staphylococcus aureus 2396 S1M10000024E08 Staphylococcus aureus 2397 S1M10000024F03 Staphylococcus aureus 2398 S1M10000024F05 Staphylococcus aureus 2400 <td></td> <td>I</td> <td>1 -2 -</td>		I	1 -2 -
2381 SIM10000024B08 Staphylococcus aureus 2382 SIM10000024B09 Staphylococcus aureus 2383 SIM10000024C02 Staphylococcus aureus 2384 SIM10000024C04 Staphylococcus aureus 2385 SIM10000024C07 Staphylococcus aureus 2386 SIM10000024D02 Staphylococcus aureus 2387 SIM10000024D03 Staphylococcus aureus 2388 SIM10000024D03 Staphylococcus aureus 2390 SIM10000024D11 Staphylococcus aureus 2391 SIM10000024E03 Staphylococcus aureus 2392 SIM10000024E03 Staphylococcus aureus 2393 SIM10000024E06 Staphylococcus aureus 2394 SIM10000024E06 Staphylococcus aureus 2395 SIM10000024E08 Staphylococcus aureus 2396 SIM10000024F02 Staphylococcus aureus 2397 SIM10000024F03 Staphylococcus aureus 2398 SIM10000024F08 Staphylococcus aureus 2400 SIM10000024G05 Staphylococcus aureus 2401 <td>1</td> <td></td> <td></td>	1		
2382 \$\text{SIM10000024B09}\$ \$\text{Staphylococcus aureus}\$ 2383 \$\text{SIM10000024C02}\$ \$\text{Staphylococcus aureus}\$ 2384 \$\text{SIM10000024C04}\$ \$\text{Staphylococcus aureus}\$ 2385 \$\text{SIM10000024C07}\$ \$\text{Staphylococcus aureus}\$ 2386 \$\text{SIM10000024D02}\$ \$\text{Staphylococcus aureus}\$ 2387 \$\text{SIM10000024D02}\$ \$\text{Staphylococcus aureus}\$ 2388 \$\text{SIM10000024D03}\$ \$\text{Staphylococcus aureus}\$ 2389 \$\text{SIM10000024D10}\$ \$\text{Staphylococcus aureus}\$ 2390 \$\text{SIM10000024E03}\$ \$\text{Staphylococcus aureus}\$ 2391 \$\text{SIM10000024E03}\$ \$\text{Staphylococcus aureus}\$ 2392 \$\text{SIM10000024E05}\$ \$\text{Staphylococcus aureus}\$ 2393 \$\text{SIM10000024E06}\$ \$\text{Staphylococcus aureus}\$ 2394 \$\text{SIM10000024E07}\$ \$\text{Staphylococcus aureus}\$ 2395 \$\text{SIM10000024F02}\$ \$\text{Staphylococcus aureus}\$ 2397 \$\text{SIM10000024F03}\$ \$\text{Staphylococcus aureus}\$ 2398 \$\text{SIM10000024F08}\$ \$Sta			
2383 \$\text{SIM10000024B10}\$ \$\text{Staphylococcus aureus}\$ 2384 \$\text{SIM10000024C02}\$ \$\text{Staphylococcus aureus}\$ 2385 \$\text{SIM10000024C07}\$ \$\text{Staphylococcus aureus}\$ 2386 \$\text{SIM10000024D02}\$ \$\text{Staphylococcus aureus}\$ 2387 \$\text{SIM10000024D03}\$ \$\text{Staphylococcus aureus}\$ 2388 \$\text{SIM10000024D10}\$ \$\text{Staphylococcus aureus}\$ 2389 \$\text{SIM10000024E01}\$ \$\text{Staphylococcus aureus}\$ 2390 \$\text{SIM10000024E03}\$ \$\text{Staphylococcus aureus}\$ 2391 \$\text{SIM10000024E05}\$ \$\text{Staphylococcus aureus}\$ 2392 \$\text{SIM10000024E05}\$ \$\text{Staphylococcus aureus}\$ 2393 \$\text{SIM10000024E06}\$ \$\text{Staphylococcus aureus}\$ 2394 \$\text{SIM10000024E07}\$ \$\text{Staphylococcus aureus}\$ 2395 \$\text{SIM10000024F02}\$ \$\text{Staphylococcus aureus}\$ 2396 \$\text{SIM10000024F03}\$ \$\text{Staphylococcus aureus}\$ 2398 \$\text{SIM10000024F08}\$ \$\text{Staphylococcus aureus}\$ 2400 \$\text{SIM10000024F08}\$ \$Sta			l
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2387 \$1M1000024D02 \$Staphylococcus aureus 2388 \$1M1000024D03 \$Staphylococcus aureus 2389 \$1M1000024D10 \$Staphylococcus aureus 2390 \$1M1000024E03 \$Staphylococcus aureus 2391 \$1M1000024E03 \$Staphylococcus aureus 2392 \$1M1000024E05 \$Staphylococcus aureus 2393 \$\$1M1000024E06 \$Staphylococcus aureus 2394 \$\$1M1000024E07 \$Staphylococcus aureus 2395 \$\$1M1000024E08 \$Staphylococcus aureus 2396 \$\$1M1000024F02 \$Staphylococcus aureus 2397 \$\$1M1000024F03 \$Staphylococcus aureus 2398 \$\$1M1000024F05 \$Staphylococcus aureus 2399 \$\$1M1000024F08 \$Staphylococcus aureus 2400 \$\$1M1000024F08 \$Staphylococcus aureus 2401 \$\$1M1000024G05 \$Staphylococcus aureus 2402 \$\$1M1000024G06 \$Staphylococcus aureus 2403 \$\$1M1000024G08 \$Staphylococcus aureus 2404 \$\$1M10000024G08 \$Staphylococcus aureus	2386	S1M10000024C07	
2389 \$1M1000024D10 Staphylococcus aureus 2390 \$1M1000024D11 Staphylococcus aureus 2391 \$1M10000024E03 Staphylococcus aureus 2392 \$1M10000024E05 Staphylococcus aureus 2393 \$1M10000024E06 Staphylococcus aureus 2394 \$1M10000024E07 Staphylococcus aureus 2395 \$1M10000024E08 Staphylococcus aureus 2396 \$51M10000024F02 Staphylococcus aureus 2397 \$1M10000024F03 Staphylococcus aureus 2398 \$1M10000024F05 Staphylococcus aureus 2399 \$1M10000024F08 Staphylococcus aureus 2400 \$1M10000024F10 Staphylococcus aureus 2401 \$1M10000024G05 Staphylococcus aureus 2402 \$1M10000024G06 Staphylococcus aureus 2403 \$1M10000024G07 Staphylococcus aureus 2404 \$1M10000024G08 Staphylococcus aureus 2405 \$1M10000024G10 Staphylococcus aureus 2406 \$1M10000024G12 Staphylococcus aureus	2387	S1M10000024D02	Staphylococcus aureus
2390 S1M1000024D11 Staphylococcus aureus 2391 S1M10000024E03 Staphylococcus aureus 2392 S1M10000024E05 Staphylococcus aureus 2393 S1M10000024E06 Staphylococcus aureus 2394 S1M10000024E07 Staphylococcus aureus 2395 S1M10000024E08 Staphylococcus aureus 2396 S1M10000024F02 Staphylococcus aureus 2397 S1M10000024F03 Staphylococcus aureus 2398 S1M10000024F05 Staphylococcus aureus 2399 S1M10000024F08 Staphylococcus aureus 2400 S1M10000024F10 Staphylococcus aureus 2401 S1M10000024G05 Staphylococcus aureus 2402 S1M10000024G06 Staphylococcus aureus 2403 S1M10000024G07 Staphylococcus aureus 2404 S1M10000024G08 Staphylococcus aureus 2405 S1M10000024G10 Staphylococcus aureus 2406 S1M10000024G12 Staphylococcus aureus	2388	S1M10000024D03	Staphylococcus aureus
2391 \$\text{SIM10000024E03}\$ \$\text{Staphylococcus aureus}\$ 2392 \$\text{SIM10000024E05}\$ \$\text{Staphylococcus aureus}\$ 2393 \$\text{SIM10000024E06}\$ \$\text{Staphylococcus aureus}\$ 2394 \$\text{SIM10000024E07}\$ \$\text{Staphylococcus aureus}\$ 2395 \$\text{SIM10000024E08}\$ \$\text{Staphylococcus aureus}\$ 2396 \$\text{SIM10000024F02}\$ \$\text{Staphylococcus aureus}\$ 2397 \$\text{SIM10000024F03}\$ \$\text{Staphylococcus aureus}\$ 2398 \$\text{SIM10000024F05}\$ \$\text{Staphylococcus aureus}\$ 2400 \$\text{SIM10000024F08}\$ \$\text{Staphylococcus aureus}\$ 2401 \$\text{SIM10000024G05}\$ \$\text{Staphylococcus aureus}\$ 2402 \$\text{SIM10000024G06}\$ \$\text{Staphylococcus aureus}\$ 2403 \$\text{SIM10000024G07}\$ \$\text{Staphylococcus aureus}\$ 2404 \$\text{SIM10000024G08}\$ \$\text{Staphylococcus aureus}\$ 2405 \$\text{SIM10000024G10}\$ \$\text{Staphylococcus aureus}\$ 2406 \$\text{SIM10000024G12}\$ \$\text{Staphylococcus aureus}\$	2389	S1M10000024D10	Staphylococcus aureus
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2394 \$\text{S1M10000024E07} \$\text{Staphylococcus aureus}\$ 2395 \$\text{S1M10000024E08} \$\text{Staphylococcus aureus}\$ 2396 \$\text{S1M10000024F02} \$\text{Staphylococcus aureus}\$ 2397 \$\text{S1M10000024F03} \$\text{Staphylococcus aureus}\$ 2398 \$\text{S1M10000024F05} \$\text{Staphylococcus aureus}\$ 2400 \$\text{S1M10000024F10} \$\text{Staphylococcus aureus}\$ 2401 \$\text{S1M10000024G05} \$\text{Staphylococcus aureus}\$ 2402 \$\text{S1M10000024G06} \$\text{Staphylococcus aureus}\$ 2403 \$\text{S1M10000024G07} \$\text{Staphylococcus aureus}\$ 2404 \$\text{S1M10000024G08} \$\text{Staphylococcus aureus}\$ 2405 \$\text{S1M10000024G10} \$\text{Staphylococcus aureus}\$ 2406 \$\text{S1M10000024G12} \$\text{Staphylococcus aureus}\$	2392	S1M10000024E05	Staphylococcus aureus
2395 S1M1000024E08 Staphylococcus aureus 2396 S1M10000024F02 Staphylococcus aureus 2397 S1M10000024F03 Staphylococcus aureus 2398 S1M10000024F05 Staphylococcus aureus 2399 S1M10000024F08 Staphylococcus aureus 2400 S1M10000024F10 Staphylococcus aureus 2401 S1M10000024G05 Staphylococcus aureus 2402 S1M10000024G06 Staphylococcus aureus 2403 S1M10000024G07 Staphylococcus aureus 2404 S1M10000024G08 Staphylococcus aureus 2405 S1M10000024G10 Staphylococcus aureus 2406 S1M10000024G12 Staphylococcus aureus	2393	S1M10000024E06	Staphylococcus aureus
2396 S1M1000024F02 Staphylococcus aureus 2397 S1M10000024F03 Staphylococcus aureus 2398 S1M10000024F05 Staphylococcus aureus 2399 S1M10000024F08 Staphylococcus aureus 2400 S1M10000024F10 Staphylococcus aureus 2401 S1M10000024G05 Staphylococcus aureus 2402 S1M10000024G06 Staphylococcus aureus 2403 S1M10000024G07 Staphylococcus aureus 2404 S1M10000024G08 Staphylococcus aureus 2405 S1M10000024G10 Staphylococcus aureus 2406 S1M10000024G12 Staphylococcus aureus	2394	S1M10000024E07	Staphylococcus aureus
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2398 S1M1000024F05 Staphylococcus aureus 2399 S1M10000024F08 Staphylococcus aureus 2400 S1M10000024F10 Staphylococcus aureus 2401 S1M10000024G05 Staphylococcus aureus 2402 S1M10000024G06 Staphylococcus aureus 2403 S1M10000024G07 Staphylococcus aureus 2404 S1M10000024G08 Staphylococcus aureus 2405 S1M10000024G10 Staphylococcus aureus 2406 S1M10000024G12 Staphylococcus aureus	2396	S1M10000024F02	Staphylococcus aureus
2399 S1M1000024F08 Staphylococus aureus 2400 S1M10000024F10 Staphylococus aureus 2401 S1M10000024G05 Staphylococus aureus 2402 S1M10000024G06 Staphylococus aureus 2403 S1M10000024G07 Staphylococus aureus 2404 S1M10000024G08 Staphylococus aureus 2405 S1M10000024G10 Staphylococus aureus 2406 S1M10000024G12 Staphylococus aureus	2397	S1M10000024F03	Staphylococcus aureus
2400 S1M1000024F10 Staphylococus aureus 2401 S1M1000024G05 Staphylococcus aureus 2402 S1M1000024G06 Staphylococcus aureus 2403 S1M1000024G07 Staphylococcus aureus 2404 S1M1000024G08 Staphylococcus aureus 2405 S1M1000024G10 Staphylococcus aureus 2406 S1M10000024G12 Staphylococcus aureus	2398	S1M10000024F05	Staphylococcus aureus
2401 SIM1000024G05 Staphylococcus aureus 2402 SIM10000024G06 Staphylococcus aureus 2403 SIM10000024G07 Staphylococcus aureus 2404 SIM10000024G08 Staphylococcus aureus 2405 SIM10000024G10 Staphylococcus aureus 2406 SIM10000024G12 Staphylococcus aureus	2399	S1M10000024F08	Staphylococcus aureus
2402 \$\text{S1M10000024G06}\$ \$\text{Staphylococcus aureus}\$ 2403 \$\text{S1M10000024G07}\$ \$\text{Staphylococcus aureus}\$ 2404 \$\text{S1M10000024G08}\$ \$\text{Staphylococcus aureus}\$ 2405 \$\text{S1M10000024G10}\$ \$\text{Staphylococcus aureus}\$ 2406 \$\text{S1M10000024G12}\$ \$\text{Staphylococcus aureus}\$	2400	S1M10000024F10	Staphylococcus aureus
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2404 S1M10000024G08 Staphylococcus aureus 2405 S1M10000024G10 Staphylococcus aureus 2406 S1M10000024G12 Staphylococcus aureus	2402	S1M10000024G06	Staphylococcus aureus
2405 S1M10000024G10 Staphylococcus aureus 2406 S1M10000024G12 Staphylococcus aureus	2403	S1M10000024G07	Staphylococcus aureus
2406 S1M10000024G12 Staphylococcus aureus		S1M10000024G08	Staphylococcus aureus
	2405	S1M10000024G10	Staphylococcus aureus
2407 S1M10000024H02 Staphylococcus aureus			Staphylococcus aureus
	2407	S1M10000024H02	Staphylococcus aureus

SeqID	Clone name	Organism
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2408	S1M1000024H04	<u> </u>
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2410	S1M1000024H08	1 2 7
2411	S1M1000025A03	Staphylococcus aureus
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2416	S1M10000025B02	Staphylococcus aureus
2417	S1M10000025B03	Staphylococcus aureus
2418	S1M10000025B05	Staphylococcus aureus
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	S1M10000025B09	Staphylococcus aureus
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2422	S1M10000025C01	Staphylococcus aureus
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2426	S1M10000025C10	Staphylococcus aureus
	S1M10000025C11	Staphylococcus aureus
2428	S1M10000025D01	Staphylococcus aureus
2429	S1M10000025D03	Staphylococcus aureus
2430	S1M10000025D04	Staphylococcus aureus
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	S1M10000025F10	Staphylococcus aureus
2444	S1M10000025F12	Staphylococcus aureus
	S1M10000025G04	Staphylococcus aureus
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2448	S1M10000025H05	Staphylococcus aureus
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2453	S1M10000026A04	Staphylococcus aureus
2454	S1M1000026A05	Staphylococcus aureus
2455	S1M10000026A06	Staphylococcus aureus
2456	S1M10000026A07	Staphylococcus aureus

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L	SeqID	Clone name	Organism
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L	2458	S1M10000026A09	Staphylococcus aureus
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	2460	S1M10000026A11	Staphylococcus aureus
ſ	2461	S1M10000026B02	Staphylococcus aureus
	2462	S1M10000026B03	Staphylococcus aureus
	2463	S1M10000026B05	Staphylococcus aureus
	2464	S1M10000026B06	Staphylococcus aureus
	2465	S1M10000026B07	Staphylococcus aureus
Ĺ	2466	S1M10000026B10	Staphylococcus aureus
Γ	2467	S1M10000026B11	Staphylococcus aureus
Γ	2468	S1M10000026B12	Staphylococcus aureus
Γ	2469	S1M10000026C01	Staphylococcus aureus
Γ	2470	S1M10000026C06	Staphylococcus aureus
Γ	2471	S1M10000026C07	Staphylococcus aureus
ľ	2472	S1M10000026C08	Staphylococcus aureus
Γ	2473	S1M10000026C11	Staphylococcus aureus
Γ	2474	S1M10000026C12	Staphylococcus aureus
Γ	2475	S1M10000026D04	Staphylococcus aureus
ľ	2476	S1M10000026D05	Staphylococcus aureus
ľ	2477	S1M10000026D06	Staphylococcus aureus
ľ	2478	S1M10000026D07	Staphylococcus aureus
Ţ	2479	S1M10000026D08	Staphylococcus aureus
Ţ	2480	S1M10000026D10	Staphylococcus aureus
ľ	2481	S1M10000026D12	Staphylococcus aureus
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ľ	2483	S1M10000026E07	Staphylococcus aureus
ľ	2484	S1M10000026E09	Staphylococcus aureus
Ī	2485	S1M10000026E10	Staphylococcus aureus
ſ	2486	S1M10000026E11	Staphylococcus aureus
ľ	2487	S1M10000026E12	Staphylococcus aureus
T	2488	S1M10000026F01	Staphylococcus aureus
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r	2500	S1M10000026G03	Staphylococcus aureus
r	2501	S1M10000026G04	Staphylococcus aureus
t	2502	S1M10000026G05	Staphylococcus aureus
r	2503	S1M10000026G06	Staphylococcus aureus
r	2504	S1M10000026G07	Staphylococcus aureus
r	2505	S1M10000026G09	Staphylococcus aureus

SeqID	Clone name	Organism
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2512	S1M10000026H05	Staphylococcus aureus
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2516	S1M10000027A04	Staphylococcus aureus
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2525	S1M10000027B11	Staphylococcus aureus
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2541	S1M10000027E03	Staphylococcus aureus Staphylococcus aureus
2542	S1M10000027E00	Staphylococcus aureus Staphylococcus aureus
2543	S1M10000027E07	Staphylococcus aureus
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2551	S1M1000027100	Staphylococcus aureus
2552	S1M1000027103	Staphylococcus aureus
2553	S1M1000027G04	Staphylococcus aureus
2554	S1M1000027G05	Staphylococcus aureus
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SeqID	Clone name	Organism
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2556	S1M1000027G07	Staphylococcus aureus
2557	S1M10000027G09	Staphylococcus aureus
2558	S1M10000027G11	Staphylococcus aureus
2559	S1M10000027H02	Staphylococcus aureus
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2564	S1M10000027H08	Staphylococcus aureus
2565	S1M10000027H09	Staphylococcus aureus
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2581	S1M10000028C04	Staphylococcus aureus
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2587	S1M10000028D04	Staphylococcus aureus
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2589	S1M1000028D07	Staphylococcus aureus
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2593	S1M1000028E08	Staphylococcus aureus
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2603	S1M10000028G04	Staphylococcus aureus
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SeqID	Clone name	Organism
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2605	S1M10000028G06	Staphylococcus aureus
2606	S1M10000028G08	Staphylococcus aureus
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2609	S1M10000028H05	Staphylococcus aureus
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2612	S1M10000029A09	Staphylococcus aureus
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2622	S1M10000029B10	Staphylococcus aureus
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2627	S1M10000029C09	Staphylococcus aureus
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2632	S1M10000029D09	Staphylococcus aureus
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2639	S1M10000029F01	Staphylococcus aureus
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2642	S1M10000029F09 S1M10000029F10	Staphylococcus aureus
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2644	S1M10000029F11 S1M10000029F12	Staphylococcus aureus
2645	S1M10000029F12 S1M10000029G01	Staphylococcus aureus
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2648	S1M1000029G02 S1M10000029G03	Staphylococcus aureus
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2650	S1M10000029G03	Staphylococcus aureus
2651	S1M1000029G07	Staphylococcus aureus
2652	S1M1000029G08	Staphylococcus aureus Staphylococcus aureus
2032	D1M10000029G1Z	Siapnyrococcus aureus

SeqID	Clone name	Organism
2653	S1M10000029H01	Staphylococcus aureus
2654	S1M10000029H05	Staphylococcus aureus
2655	S1M10000029H06	Staphylococcus aureus
2656	S1M10000029H08	Staphylococcus aureus
2657	S1M10000029H09	Staphylococcus aureus
2658	S1M10000029H10	Staphylococcus aureus
2659	S1M10000030A02	Staphylococcus aureus
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2661	S1M10000030A09	Staphylococcus aureus
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2665	S1M10000030B05	Staphylococcus aureus
2666	S1M10000030B07	Staphylococcus aureus
2667	S1M10000030B09	Staphylococcus aureus
2668	S1M10000030C02	Staphylococcus aureus
2669	S1M10000030C03	Staphylococcus aureus
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2671	S1M10000030C05	Staphylococcus aureus
2672	S1M10000030C08	Staphylococcus aureus
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2682	S1M10000030D09	Staphylococcus aureus
2683	S1M10000030D10	Staphylococcus aureus
2684	S1M10000030D11	Staphylococcus aureus
2685	S1M10000030E02	Staphylococcus aureus
2686	S1M10000030E06	Staphylococcus aureus
2687	S1M10000030E07	Staphylococcus aureus
2688	S1M10000030E11	Staphylococcus aureus
2689	S1M10000030E12	Staphylococcus aureus
2690	S1M10000030F01	Staphylococcus aureus
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2696	S1M10000030G05	Staphylococcus aureus
2697	S1M10000030G07	Staphylococcus aureus
2698	S1M10000030G08	Staphylococcus aureus
2699	S1M10000030G09	Staphylococcus aureus
2700	S1M10000030G10	Staphylococcus aureus
2701	S1M10000030G11	Staphylococcus aureus

770333	IABI	SE IA	1 € 1/6501/02100
SeqID	Clone name	Organism	
2702	S1M10000030G12	Staphylococcus aureus	
2703	S1M10000030H01	Staphylococcus aureus	· · · · · · · · · · · · · · · · · · ·
2704	S1M10000030H02	Staphylococcus aureus	
2705	S1M10000030H03	Staphylococcus aureus	
2706	S1M10000030H05	Staphylococcus aureus	
2707	S1M10000030H07	Staphylococcus aureus	
2708	S1M10000030H09	Staphylococcus aureus	
2709	S1M10000031A03	Staphylococcus aureus	
2710	S1M10000031A08	Staphylococcus aureus	· · · · · · · · · · · · · · · · · · ·
2711	S1M10000031A10	Staphylococcus aureus	
2712	SIM10000031B01	Staphylococcus aureus	
2713	S1M10000031B02	Staphylococcus aureus	
2714	S1M10000031B04	Staphylococcus aureus	
2715	S1M10000031B11	Staphylococcus aureus	
2716	S1M10000031B12	Staphylococcus aureus	
2717	S1M10000031C04	Staphylococcus aureus	
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2719	S1M10000031C09	Staphylococcus aureus	
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2728	S1M10000031E07	Staphylococcus aureus	
2729	\$1M10000031E08	Staphylococcus aureus	
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2739	S1M10000031F12	Staphylococcus aureus	
2740	S1M10000031G02	Staphylococcus aureus	
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2742	S1M10000031G04	Staphylococcus aureus	
2743	S1M10000031G06	Staphylococcus aureus	
2744	S1M10000031G09	Staphylococcus aureus	
2745	S1M10000031G10	Staphylococcus aureus	
2746	S1M10000031G11	Staphylococcus aureus	
2747	S1M10000031H01	Staphylococcus aureus	
2748	S1M10000031H02	Staphylococcus aureus	
2749	S1M10000031H06	Staphylococcus aureus	
2750	S1M10000031H09	Staphylococcus aureus	

SeqID	Clone name	Organism
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2752	S1M10000032A03	Staphylococcus aureus
2753	S1M10000032A05	Staphylococcus aureus
2754	S1M10000032A06	Staphylococcus aureus
2755	S1M10000032A07	Staphylococcus aureus
2756	S1M10000032A08	Staphylococcus aureus
2757	S1M10000032A10	Staphylococcus aureus
2758	S1M10000032B01	Staphylococcus aureus
2759	S1M10000032B05	Staphylococcus aureus
2760	S1M10000032B07	Staphylococcus aureus
2761	S1M10000032B08	Staphylococcus aureus
2762	S1M10000032B11	Staphylococcus aureus
2763	S1M10000032B12	Staphylococcus aureus
2764	S1M10000032C01	Staphylococcus aureus
2765	S1M10000032C03	Staphylococcus aureus
2766	S1M10000032C04	Staphylococcus aureus
2767	S1M10000032C05	Staphylococcus aureus
2768	S1M10000032C09	Staphylococcus aureus
2769	S1M10000032C10	Staphylococcus aureus
2770	S1M10000032C11	Staphylococcus aureus
2771	S1M10000032C12	Staphylococcus aureus
2772	S1M10000032D03	Staphylococcus aureus
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2774	S1M10000032D07	Staphylococcus aureus
2775	S1M10000032D09	Staphylococcus aureus
2776	S1M10000032D11 S1M10000032E02	Staphylococcus aureus
2778	S1M10000032E02	Staphylococcus aureus Staphylococcus aureus
2779	S1M10000032E03	Staphylococcus aureus Staphylococcus aureus
2780	S1M10000032E04	Staphylococcus aureus
2781	S1M1000032E08	Staphylococcus aureus
2782	S1M10000032E09	Staphylococcus aureus
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2791	S1M10000032F12	Staphylococcus aureus
2792	S1M10000032G02	Staphylococcus aureus
2793	S1M10000032G03	Staphylococcus aureus
2794	S1M10000032G04	Staphylococcus aureus
2795	S1M10000032G06	Staphylococcus aureus
2796	S1M10000032G08	Staphylococcus aureus
2797	S1M10000032G10	Staphylococcus aureus
2798	S1M10000032G12	Staphylococcus aureus
2799	S1M10000032H01	Staphylococcus aureus

SeqID	Clone name	Organism
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2805	S1M10000033A07	Staphylococcus aureus
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2807	S1M10000033A10	Staphylococcus aureus
2808	S1M10000033B02	Staphylococcus aureus
2809	S1M10000033B07	Staphylococcus aureus
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2814	S1M10000033D02	Staphylococcus aureus
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2829	S1M10000033F11	Staphylococcus aureus
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2838	S1M10000033H03	Staphylococcus aureus
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2845	S1M10000034A03 S1M10000034A04	Staphylococcus aureus
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2848	911V110000034A08	Staphylococcus aureus

SeqID	Clone name	Organism
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2862	S1M10000034C07	Staphylococcus aureus
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2865	S1M10000034D01	Staphylococcus aureus
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2867	S1M10000034D06	Staphylococcus aureus
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2885	S1M10000034F04	Staphylococcus aureus
2886	S1M10000034F05	Staphylococcus aureus
2887	S1M10000034F07	Staphylococcus aureus
2888	S1M1000034F08	Staphylococcus aureus
2889	S1M1000034F09	Staphylococcus aureus
2890	S1M10000034F10	Staphylococcus aureus
2891	S1M10000034F12	Staphylococcus aureus
2892	S1M10000034G02	Staphylococcus aureus
2893	S1M10000034G03	Staphylococcus aureus
2894 2895	S1M10000034G06 S1M10000034G07	Staphylococcus aureus
		Staphylococcus aureus
2896	S1M1000034G08	Staphylococcus aureus
2897	S1M10000034G09	Staphylococcus aureus

2898 S1M10000034G11 Staphylococcus aureus 2899 S1M10000034G12 Staphylococcus aureus 2900 S1M10000034H01 Staphylococcus aureus 2901 S1M10000034H02 Staphylococcus aureus	
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2901 S1M10000034H02 Staphylococcus aureus	
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2903 S1M10000034H06 Staphylococcus aureus	
2904 S1M10000034H07 Staphylococcus aureus	
2905 S1M10000034H08 Staphylococcus aureus	
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2908 S1M10000035A03 Staphylococcus aureus	
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2918 S1M10000035B11 Staphylococcus aureus	
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2920 S1M10000035C02 Staphylococcus aureus	
2921 S1M10000035C04 Staphylococcus aureus	
2922 \$1M10000035C06	
2923 \$1M1000035C11 \$Staphylococcus aureus 2924 \$1M1000035D01 \$Staphylococcus aureus	
2926 S1M10000035D06 Staphylococcus aureus 2927 S1M10000035D09 Staphylococcus aureus	
2928 S1M10000035D12 Staphylococcus aureus	
2929 S1M10000035E02 Staphylococcus aureus	
2930 S1M1000035E03 Staphylococcus aureus	
2931 S1M1000035E04 Staphylococcus aureus	
2932 S1M1000035E08 Staphylococcus aureus	
2933 S1M10000035E09 Staphylococcus aureus	
2934 S1M10000035E12 Staphylococcus aureus	
2935 S1M10000035F03 Staphylococcus aureus	- -
2936 S1M10000035F04 Staphylococcus aureus	
2937 S1M10000035F09 Staphylococcus aureus	
2938 S1M10000035F12 Staphylococcus aureus	
2939 S1M10000035G02 Staphylococcus aureus	
2940 S1M10000035G09 Staphylococcus aureus	
2941 S1M10000035G11 Staphylococcus aureus	
2942 S1M10000035G12 Staphylococcus aureus	
2943 S1M10000035H01 Staphylococcus aureus	
2944 S1M10000035H07 Staphylococcus aureus	
2945 S1M10000035H08 Staphylococcus aureus	
2946 S1M10000035H09 Staphylococcus aureus	

SeqID	Clone name	Organism
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2948	S1M10000035H11	Staphylococcus aureus
2949	S1M10000036A02	Staphylococcus aureus
2950	S1M10000036A03	Staphylococcus aureus
2951	SIM10000036A04	Staphylococcus aureus
2952	S1M10000036A05	Staphylococcus aureus
2953	S1M10000036A08	Staphylococcus aureus
2954	S1M10000036A11	Staphylococcus aureus
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2956	S1M10000036B04	Staphylococcus aureus
2957	S1M10000036B06	Staphylococcus aureus
2958	S1M10000036B07	Staphylococcus aureus
2959	S1M10000036B08	Staphylococcus aureus
2960	S1M10000036B11	Staphylococcus aureus
2961	S1M10000036B12	Staphylococcus aureus
2962	S1M10000036C01	Staphylococcus aureus
2963	S1M10000036C03	Staphylococcus aureus
2964	S1M10000036C04	Staphylococcus aureus
2965	S1M10000036C05	Staphylococcus aureus
2966	S1M10000036C06	Staphylococcus aureus
2967	S1M10000036C07	Staphylococcus aureus
2968	S1M10000036C09	Staphylococcus aureus
2969	S1M10000036C10	Staphylococcus aureus
2970	S1M10000036D02	Staphylococcus aureus
2971	S1M10000036D03	Staphylococcus aureus
2972	S1M10000036D06	Staphylococcus aureus
2973	S1M10000036D08	Staphylococcus aureus
2974	S1M10000036D10	Staphylococcus aureus
2975	S1M10000036D11	Staphylococcus aureus
2976	S1M10000036D12	Staphylococcus aureus
2977	S1M10000036E06	Staphylococcus aureus
2978	S1M10000036E08	Staphylococcus aureus
2979	S1M10000036E11	Staphylococcus aureus
2980	S1M10000036F06	Staphylococcus aureus
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2982	S1M1000036F08	Staphylococcus aureus
2983	S1M10000036F09	Staphylococcus aureus
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2985	S1M10000036F11	Staphylococcus aureus
2986	S1M1000036G03	Staphylococcus aureus
2987 2988	S1M10000036G07	Staphylococcus aureus
2988	S1M10000036G08 S1M10000036G11	Staphylococcus aureus
2989		Staphylococcus aureus
2990	S1M10000036H01 S1M10000036H02	Staphylococcus aureus
2991	S1M10000036H02	Staphylococcus aureus
2992	S1M10000036H03	Staphylococcus aureus
2993	S1M10000036H04	Staphylococcus aureus
2994	S1M10000036H06	Staphylococcus aureus
2993	STMITOOOOSOMOO	Staphylococcus aureus

SeqID	Clone name	Organism
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2997	S1M10000036H11	Staphylococcus aureus
2998	S1M10000037A02	Staphylococcus aureus
2999	S1M10000037A03	Staphylococcus aureus
3000	S1M10000037A06	Staphylococcus aureus
3001	S1M10000037A08	Staphylococcus aureus
3002	S1M10000037A09	Staphylococcus aureus
3003	SIM10000037A11	Staphylococcus aureus
3004	S1M10000037A12	Staphylococcus aureus
3005	S1M10000037B03	Staphylococcus aureus
3006	S1M10000037B04	Staphylococcus aureus
3007	S1M10000037B05	Staphylococcus aureus
3008	S1M10000037B06	Staphylococcus aureus
3009	S1M10000037B07	Staphylococcus aureus
3010	S1M10000037B08	Staphylococcus aureus
3011	S1M10000037B10	Staphylococcus aureus
3012	S1M10000037B11	Staphylococcus aureus
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3018	S1M10000037C09	Staphylococcus aureus
3019	S1M10000037C10	Staphylococcus aureus
3020	S1M10000037D04	Staphylococcus aureus
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3023	S1M10000037D09	Staphylococcus aureus
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3027	S1M10000037E08	Staphylococcus aureus
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3030	S1M10000037E10	Staphylococcus aureus Staphylococcus aureus
3031	S1M1000037E11	Staphylococcus aureus
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3034	S1M1000037F02	Staphylococcus aureus
3035	S1M1000037F04	Staphylococcus aureus
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3040	S1M10000037F09	Staphylococcus aureus
3041	S1M1000037F10	Staphylococcus aureus
3042	S1M1000037F10	Staphylococcus aureus
3043	S1M1000037G01	Staphylococcus aureus
3044	S1M1000037G02	Staphylococcus aureus
	5211110000057005	Dispreysocooms amens

SeqID	Clone name	Organism
3045	S1M10000037G06	Staphylococcus aureus
3046	S1M10000037G07	Staphylococcus aureus
3047	S1M10000037G08	Staphylococcus aureus
3048	S1M10000037G10	Staphylococcus aureus
3049	S1M10000037H02	Staphylococcus aureus
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3062	S1M10000038B01	Staphylococcus aureus
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3078	S1M10000038D08 S1M10000038D09	Staphylococcus aureus
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3082	S1M1000038D12 S1M10000038E01	Staphylococcus aureus
3084	S1M1000038E01	Staphylococcus aureus Staphylococcus aureus
3085	S1M10000038E02	Staphylococcus aureus .
3086	S1M1000038E03	Staphylococcus aureus . Staphylococcus aureus
3087	S1M1000038E04	Staphylococcus aureus Staphylococcus aureus
3088	S1M1000038E05	Staphylococcus aureus Staphylococcus aureus
3089	S1M1000038E07	Staphylococcus aureus Staphylococcus aureus
3090	S1M1000038E07	Staphylococcus aureus
3090	S1M1000038E10	Staphylococcus aureus Staphylococcus aureus
3092	S1M1000038E12	Staphylococcus aureus Staphylococcus aureus
3092	S1M1000038F03	Staphylococcus aureus Staphylococcus aureus
3093	D1M10000030F04	Siapnyiococcus aureus

SeqID	Clone name	Organism
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3095	S1M10000038F06	Staphylococcus aureus
3096	S1M10000038F08	Staphylococcus aureus
3097	S1M10000038F09	Staphylococcus aureus
3098	S1M10000038F10	Staphylococcus aureus
3099	S1M10000038F11	Staphylococcus aureus
3100	S1M10000038F12	Staphylococcus aureus
3101	S1M10000038G01	Staphylococcus aureus
3102	S1M10000038G03	Staphylococcus aureus
3103	S1M10000038G04	Staphylococcus aureus
3104	S1M10000038G06	Staphylococcus aureus
3105	S1M10000038G08	Staphylococcus aureus
3106	S1M10000038G10	Staphylococcus aureus
3107	S1M10000038G11	Staphylococcus aureus
3108	S1M10000038G12	Staphylococcus aureus
3109	S1M10000038H03	Staphylococcus aureus
3110	S1M10000038H07	Staphylococcus aureus
3111	S1M10000038H09	Staphylococcus aureus
3112	S1M10000038H11	Staphylococcus aureus
3113	S1M10000039A02	Staphylococcus aureus
3114	S1M10000039A05	Staphylococcus aureus
3115	S1M10000039A07	Staphylococcus aureus
3116	S1M10000039A08	Staphylococcus aureus
3117	S1M10000039A11	Staphylococcus aureus
3118	S1M10000039A12	Staphylococcus aureus
3119 3120	S1M10000039B02	Staphylococcus aureus
3120	S1M10000039B06 S1M10000039B07	Staphylococcus aureus
3122	S1M1000039B07	Staphylococcus aureus Staphylococcus aureus
3123	S1M1000039B10	Staphylococcus aureus
3124	S1M10000039B12	Staphylococcus aureus
3125	S1M1000039C06	Staphylococcus aureus
3126	S1M1000039C07	Staphylococcus aureus
3127	S1M10000039C08	Staphylococcus aureus
3128	S1M10000039C09	Staphylococcus aureus
3129	S1M10000039C10	Staphylococcus aureus
3130	S1M10000039C11	Staphylococcus aureus
3131	S1M10000039D02	Staphylococcus aureus
3132	S1M10000039D09	Staphylococcus aureus
3133	S1M10000039D10	Staphylococcus aureus
3134	S1M10000039E01	Staphylococcus aureus
3135	S1M10000039E08	Staphylococcus aureus
3136	S1M10000039E09	Staphylococcus aureus
3137	S1M10000039E10	Staphylococcus aureus
3138	S1M10000039E11	Staphylococcus aureus
3139	S1M10000039F02	Staphylococcus aureus
3140	S1M10000039F03	Staphylococcus aureus
3141	S1M10000039F05	Staphylococcus aureus
3142	S1M10000039F07	Staphylococcus aureus

SeqID	Clone name	Organism
3143	S1M10000039F08	Staphylococcus aureus
3144	S1M10000039F09	Staphylococcus aureus
3145	S1M10000039F10	Staphylococcus aureus
3146	S1M10000039F12	Staphylococcus aureus
3147	S1M10000039G03	Staphylococcus aureus
3148	S1M10000039G04	Staphylococcus aureus
3149	S1M10000039G07	Staphylococcus aureus
3150	S1M10000039G10	Staphylococcus aureus
3151	S1M10000039H02	Staphylococcus aureus
3152	S1M10000039H03	Staphylococcus aureus
3153	S1M10000039H04	Staphylococcus aureus
3154	S1M10000039H06	Staphylococcus aureus
3155	S1M10000039H07	Staphylococcus aureus
3156	S1M10000039H08	Staphylococcus aureus
3157	S1M10000040A04	Staphylococcus aureus
3158	S1M10000040A05	Staphylococcus aureus
3159	S1M10000040A07	Staphylococcus aureus
3160	S1M10000040A08	Staphylococcus aureus
3161	S1M10000040A10	Staphylococcus aureus
3162	S1M10000040A11	Staphylococcus aureus
3163	S1M10000040B01	Staphylococcus aureus
3164	S1M10000040B03	Staphylococcus aureus
3165	S1M10000040B07	Staphylococcus aureus
3166	S1M10000040B11	Staphylococcus aureus
3167	S1M10000040C03	Staphylococcus aureus
3168	S1M10000040C04	Staphylococcus aureus
3169	S1M10000040C05	Staphylococcus aureus
3170	S1M10000040C06	Staphylococcus aureus
3171	S1M10000040C07	Staphylococcus aureus
3172 3173	S1M10000040C08 S1M10000040C10	Staphylococcus aureus
3173	S1M10000040C10	Staphylococcus aureus
3174	S1M10000040C11	Staphylococcus aureus
3176	S1M1000040D01	Staphylococcus aureus
3177	S1M10000040D03	Staphylococcus aureus Staphylococcus aureus
3178	S1M1000040D08	
3179	S1M10000040D09	Staphylococcus aureus Staphylococcus aureus
3180	S1M1000040D11	Staphylococcus aureus Staphylococcus aureus
3181	S1M10000040E01	Staphylococcus aureus Staphylococcus aureus
3182	S1M1000040E02	Staphylococcus aureus
3183	S1M1000040E05	Staphylococcus aureus
3184	S1M1000040E05	Staphylococcus aureus
3185	S1M1000040E00	Staphylococcus aureus
3186	S1M1000040E07	Staphylococcus aureus
3187	S1M10000040E09	Staphylococcus aureus Staphylococcus aureus
3188	S1M10000040E10	Staphylococcus aureus
3189	S1M10000040E11	Staphylococcus aureus
3190	S1M1000040E01	Staphylococcus aureus
3191	S1M1000040F02	Staphylococcus aureus
	21111000010102	Diaphytococas aareas

Γ	SeqID	Clone name	Organism
ľ	3192	S1M10000040F03	Staphylococcus aureus
	3193	\$1M10000040F04	Staphylococcus aureus
1	3194	S1M10000040F05	Staphylococcus aureus
1	3195	S1M10000040F06	Staphylococcus aureus
	3196	S1M10000040F08	Staphylococcus aureus
Ī	3197	S1M10000040F09	Staphylococcus aureus
	3198	S1M10000040F12	Staphylococcus aureus
ſ	3199	S1M10000040G01	Staphylococcus aureus
F	3200	S1M10000040G02	Staphylococcus aureus
	3201	S1M10000040G04	Staphylococcus aureus
	3202	S1M10000040G07	Staphylococcus aureus
. [3203	\$1M10000040G08	Staphylococcus aureus
[3204	S1M10000040G12	Staphylococcus aureus
	3205	S1M10000040H02	Staphylococcus aureus
[3206	S1M10000040H03	Staphylococcus aureus
	3207	S1M10000040H04	Staphylococcus aureus
Ĺ	3208	S1M10000040H05	Staphylococcus aureus
	3209	S1M10000040H07	Staphylococcus aureus
Ĺ	3210	S1M10000040H10	Staphylococcus aureus
	3211	S1M10000041A03	Staphylococcus aureus
Ĺ	3212	S1M10000041B02	Staphylococcus aureus
	3213	S1M10000041B03	Staphylococcus aureus
	3214	S1M10000041B05	Staphylococcus aureus
	3215	S1M10000041B06	Staphylococcus aureus
4	3216	S1M10000041B07	Staphylococcus aureus
	3217	S1M10000041B12	Staphylococcus aureus
	3218	S1M10000041C08	Staphylococcus aureus
1	3219	S1M10000041C10	Staphylococcus aureus
<u> </u>	3220 3221	S1M10000041C11 S1M10000041D06	Staphylococcus aureus
Ļ	3222		Staphylococcus aureus
	3223	S1M10000041D07 S1M10000041D08	Staphylococcus aureus
-	3223	S1M10000041D08	Staphylococcus aureus
-	3225	S1M10000041D10	Staphylococcus aureus
-	3226	S1M10000041D12	Staphylococcus aureus Staphylococcus aureus
-	3227	S1M1000041E06	Staphylococcus aureus Staphylococcus aureus
-	3228	S1M1000041E00	Staphylococcus aureus Staphylococcus aureus
-	3229	S1M1000041E09	Staphylococcus aureus Staphylococcus aureus
-	3230	S1M1000041E12	Staphylococcus aureus Staphylococcus aureus
-	3231	S1M1000041103	Staphylococcus aureus
-	3232	S1M10000041F12	Staphylococcus aureus
-	3233	S1M10000041G01	Staphylococcus aureus
-	3234	S1M10000041G06	Staphylococcus aureus
-	3235	S1M1000041G08	Staphylococcus aureus
-	3236	S1M10000041G10	Staphylococcus aureus
-	3237	S1M1000041G11	Staphylococcus aureus
}	3238	S1M1000041H01	Staphylococcus aureus
}	3239	S1M10000041H04	Staphylococcus aureus
H	3240	S1M10000041H05	Staphylococcus aureus
L			L A a c c a a m m m m m

SeqID	Clone name	Organism
3241	S1M10000041H07	Staphylococcus aureus
3242	S1M10000041H08	Staphylococcus aureus
3243	S1M10000041H09	Staphylococcus aureus
3244	S1M10000042A04	Staphylococcus aureus
3245	S1M10000042A05	Staphylococcus aureus
3246	S1M10000042A06	Staphylococcus aureus
3247	S1M10000042A07	Staphylococcus aureus
3248	S1M10000042A09	Staphylococcus aureus
3249	S1M10000042A11	Staphylococcus aureus
3250	S1M10000042A12	Staphylococcus aureus
3251	S1M10000042B02	Staphylococcus aureus
3252	S1M10000042B03	Staphylococcus aureus
3253	S1M10000042B06	Staphylococcus aureus
3254	S1M10000042B07	Staphylococcus aureus
3255	S1M10000042B08	Staphylococcus aureus
3256	S1M10000042B09	Staphylococcus aureus
3257	S1M10000042B10	Staphylococcus aureus
3258	S1M10000042B11	Staphylococcus aureus
3259	S1M10000042B12	Staphylococcus aureus
3260	S1M10000042C02	Staphylococcus aureus
3261	S1M10000042C06	Staphylococcus aureus
3262	S1M10000042C10	Staphylococcus aureus
3263	S1M10000042C11	Staphylococcus aureus
3264	S1M10000042D04	Staphylococcus aureus
3265	S1M10000042D07	Staphylococcus aureus
3266	S1M10000042D10	Staphylococcus aureus
3267	S1M10000042D11	Staphylococcus aureus
3268	S1M10000042E03	Staphylococcus aureus
3269	S1M1000042E06	Staphylococcus aureus
3270 3271	S1M10000042E08 S1M10000042F01	Staphylococcus aureus
3271		Staphylococcus aureus
3273	S1M10000042F02 S1M10000042F05	Staphylococcus aureus
3274	S1M10000042F05 S1M10000042F06	Staphylococcus aureus
3275	S1M1000042F08	Staphylococcus aureus
3276	S1M1000042F08	Staphylococcus aureus
3277	S1M10000042F09	Staphylococcus aureus
3278	S1M10000042F10	Staphylococcus aureus
3279	S1M10000042F11 S1M10000042G01	Staphylococcus aureus Staphylococcus aureus
3280	S1M1000042G01	Staphylococcus aureus Staphylococcus aureus
3281	S1M1000042G03	Staphylococcus aureus
3282	S1M10000042G09	Staphylococcus aureus Staphylococcus aureus
3283	S1M1000042G09	Staphylococcus aureus Staphylococcus aureus
3284	S1M10000042G12	Staphylococcus aureus Staphylococcus aureus
3285	S1M10000042H07	Staphylococcus aureus
3286	S1M10000042H07	Staphylococcus aureus
3287	S1M100000421111 S1M10000043A02	Staphylococcus aureus
3288	S1M10000043A02	Staphylococcus aureus
3289	S1M10000043A03	Staphylococcus aureus
J207	511111000043104	этартугососсия интеиз

SeqID	Clone name	Organism
3290	S1M10000043A06	Staphylococcus aureus
3291	S1M10000043A07	Staphylococcus aureus
3292	S1M10000043A08	Staphylococcus aureus
3293	S1M10000043A10	Staphylococcus aureus
3294	S1M10000043A11	Staphylococcus aureus
3295	S1M10000043A12	Staphylococcus aureus
3296	S1M10000043B01	Staphylococcus aureus
3297	S1M10000043B02	Staphylococcus aureus
3298	S1M10000043B07	Staphylococcus aureus
3299	S1M10000043B08	Staphylococcus aureus
3300	S1M10000043B09	Staphylococcus aureus
3301	S1M10000043B10	Staphylococcus aureus
3302	S1M10000043B12	Staphylococcus aureus
3303	S1M10000043C02	Staphylococcus aureus
3304	S1M10000043C07	Staphylococcus aureus
3305	S1M10000043C11	Staphylococcus aureus
3306	S1M10000043C12	Staphylococcus aureus
3307	S1M10000043D01	Staphylococcus aureus
3308	S1M10000043D02	Staphylococcus aureus
3309	S1M10000043D04	Staphylococcus aureus
3310	S1M10000043D10	Staphylococcus aureus
3311	S1M10000043D12	Staphylococcus aureus
3312	S1M10000043E02	Staphylococcus aureus
3313	S1M10000043E03	Staphylococcus aureus
3314	S1M10000043E05	Staphylococcus aureus
3315	S1M10000043E07	Staphylococcus aureus
3316 3317	S1M10000043E08 S1M10000043E10	Staphylococcus aureus
3317		Staphylococcus aureus
3319	S1M10000043E11 S1M10000043E12	Staphylococcus aureus Staphylococcus aureus
3320	S1M10000043E12	Staphylococcus aureus
3321	S1M10000043F01	Staphylococcus aureus
3322	S1M1000043F07	Staphylococcus aureus
3323	S1M10000043F08	Staphylococcus aureus
3324	S1M10000043F09	Staphylococcus aureus
3325	S1M1000043G01	Staphylococcus aureus
3326	S1M10000043G04	Staphylococcus aureus
3327	S1M10000043G05	Staphylococcus aureus
3328	S1M10000043G09	Staphylococcus aureus
3329	S1M10000043G10	Staphylococcus aureus
3330	S1M10000043H01	Staphylococcus aureus
3331	S1M10000043H03	Staphylococcus aureus
3332	S1M10000043H04	Staphylococcus aureus
3333	S1M10000043H05	Staphylococcus aureus
3334	S1M10000043H06	Staphylococcus aureus
3335	S1M10000043H09	Staphylococcus aureus
3336	S1M10000043H10	Staphylococcus aureus
3337	S1M10000043H11	Staphylococcus aureus
3338	S1M10000044A02	Staphylococcus aureus

SeqID	Clone name	Organism
3339	S1M10000044A06	Staphylococcus aureus
3340	S1M10000044A08	Staphylococcus aureus
3341	S1M10000044A09	Staphylococcus aureus
3342	S1M10000044A11	Staphylococcus aureus
3343	S1M10000044A12	Staphylococcus aureus
3344	S1M10000044B01	Staphylococcus aureus
3345	S1M10000044B02	Staphylococcus aureus
3346	S1M10000044B05	Staphylococcus aureus
3347	S1M10000044B06	Staphylococcus aureus
3348	S1M10000044B08	Staphylococcus aureus
3349	S1M10000044B11	Staphylococcus aureus
3350	S1M10000044B12	Staphylococcus aureus
3351	S1M10000044C04	Staphylococcus aureus
3352	S1M10000044C06	Staphylococcus aureus
3353	S1M10000044C07	Staphylococcus aureus
3354	S1M10000044C08	Staphylococcus aureus
3355	S1M10000044C11	Staphylococcus aureus
3356	S1M10000044C12	Staphylococcus aureus
3357	S1M10000044D01	Staphylococcus aureus
3358	S1M10000044D04	Staphylococcus aureus
3359	S1M10000044D06	Staphylococcus aureus
3360	S1M10000044D08	Staphylococcus aureus
3361	S1M10000044D09	Staphylococcus aureus
3362	S1M10000044D10	Staphylococcus aureus
3363	S1M10000044D11	Staphylococcus aureus
3364	S1M10000044D12	Staphylococcus aureus
3365	S1M10000044E01	Staphylococcus aureus
3366	S1M10000044E02	Staphylococcus aureus
3367	S1M10000044E06	Staphylococcus aureus
3368	S1M10000044E07	Staphylococcus aureus
3369	S1M10000044E09	Staphylococcus aureus
3370 3371	S1M10000044E10 S1M10000044E11	Staphylococcus aureus
3371	S1M10000044E11 S1M10000044F02	Staphylococcus aureus
3373	S1M10000044F02	Staphylococcus aureus
3374	S1M10000044F08	Staphylococcus aureus
3375	S1M10000044F08	Staphylococcus aureus
3376	S1M10000044F10	Staphylococcus aureus Staphylococcus aureus
3377	S1M10000044G02	Staphylococcus aureus Staphylococcus aureus
3378	S1M1000044G08	Staphylococcus aureus Staphylococcus aureus
3379	S1M1000044G08	Staphylococcus aureus
3380	S1M1000044G10	Staphylococcus aureus
3381	S1M10000044G11	Staphylococcus aureus Staphylococcus aureus
3382	S1M10000044H00	Staphylococcus aureus Staphylococcus aureus
3383	S1M10000044H07	Staphylococcus aureus
3384	S1M1000044H08	Staphylococcus aureus Staphylococcus aureus
3385	S1M10000044H09	Staphylococcus aureus Staphylococcus aureus
3386	S1M10000044H10	Staphylococcus aureus
3387	S1M10000044H11	Staphylococcus aureus
3307	511110000043/402	Supriyiococcus aureus

SeqID	Clone name	Organism
3388	S1M10000045A06	Staphylococcus aureus
3389	S1M10000045A07	Staphylococcus aureus
3390	S1M10000045A08	Staphylococcus aureus
3391	S1M10000045A12	Staphylococcus aureus
3392	S1M10000045B01	Staphylococcus aureus
3393	S1M10000045B02	Staphylococcus aureus
3394	S1M10000045B03	Staphylococcus aureus
3395	S1M10000045B07	Staphylococcus aureus
3396	S1M10000045B10	Staphylococcus aureus
3397	S1M10000045B11	Staphylococcus aureus
3398	S1M10000045B12	Staphylococcus aureus
3399	S1M10000045C02	Staphylococcus aureus
3400	S1M10000045C03	Staphylococcus aureus
3401	S1M10000045C04	Staphylococcus aureus
3402	S1M10000045C05	Staphylococcus aureus
3403	S1M10000045C07	Staphylococcus aureus
3404	S1M10000045C09	Staphylococcus aureus
3405	S1M10000045D01	Staphylococcus aureus
3406	S1M10000045D03	Staphylococcus aureus
3407	S1M10000045D07	Staphylococcus aureus
3408	S1M10000045D08	Staphylococcus aureus
3409	S1M10000045D09	Staphylococcus aureus
3410	S1M10000045D10	Staphylococcus aureus
3411	S1M10000045D11	Staphylococcus aureus
3412	S1M10000045D12	Staphylococcus aureus
3413	S1M10000045E04	Staphylococcus aureus
3414	S1M10000045E05	Staphylococcus aureus
3415	S1M10000045E08	Staphylococcus aureus
3416	S1M10000045E09	Staphylococcus aureus
3417	S1M10000045E10	Staphylococcus aureus
3418	S1M10000045E11	Staphylococcus aureus
3419	S1M10000045E12	Staphylococcus aureus
3420	S1M10000045F04	Staphylococcus aureus
3421	S1M10000045F05	Staphylococcus aureus
3422	S1M10000045F08	Staphylococcus aureus
3423	S1M10000045F11	Staphylococcus aureus
3424	S1M10000045F12	Staphylococcus aureus
3425	S1M10000045G03	Staphylococcus aureus
3426	S1M10000045G06	Staphylococcus aureus
3427	S1M10000045G07	Staphylococcus aureus
3428	S1M10000045G08	Staphylococcus aureus
3429	S1M10000045G10	Staphylococcus aureus
3430	S1M10000045G12	Staphylococcus aureus
3431	S1M10000045H06	Staphylococcus aureus
3432	S1M10000045H10	Staphylococcus aureus
3433	S1M10000045H11	Staphylococcus aureus
3434	S1M10000046A03	Staphylococcus aureus
3435	S1M10000046A04	Staphylococcus aureus
3436	S1M10000046A06	Staphylococcus aureus

SeqID	Clone name	Organism
3437	S1M10000046A08	Staphylococcus aureus
3438	S1M10000046A09	Staphylococcus aureus
3439	S1M10000046A11	Staphylococcus aureus
3440	S1M10000046A12	Staphylococcus aureus
3441	S1M10000046B01	Staphylococcus aureus
3442	S1M10000046B03	Staphylococcus aureus
3443	S1M10000046B04	Staphylococcus aureus
3444	S1M10000046B05	Staphylococcus aureus
3445	S1M10000046B07	Staphylococcus aureus
3446	S1M10000046B08	Staphylococcus aureus
3447	S1M10000046B09	Staphylococcus aureus
3448	S1M10000046B11	Staphylococcus aureus
3449	S1M10000046B12	Staphylococcus aureus
3450	S1M10000046C02	Staphylococcus aureus
3451	S1M10000046C04	Staphylococcus aureus
3452	S1M10000046C05	Staphylococcus aureus
3453	S1M10000046C06	Staphylococcus aureus
3454	S1M10000046C07	Staphylococcus aureus
3455	S1M10000046C08	Staphylococcus aureus
3456	S1M10000046C11	Staphylococcus aureus
3457	S1M10000046C12	Staphylococcus aureus
3458	S1M10000046D01	Staphylococcus aureus
3459	S1M10000046D02	Staphylococcus aureus
3460	S1M10000046D03	Staphylococcus aureus
3461	S1M10000046D04	Staphylococcus aureus
3462	S1M1000046D05	Staphylococcus aureus
3463	S1M10000046D08	Staphylococcus aureus
3464	S1M10000046D09	Staphylococcus aureus
3465	S1M10000046D10	Staphylococcus aureus
3466	S1M10000046D11	Staphylococcus aureus
3467 3468	S1M10000046D12 S1M10000046E01	Staphylococcus aureus
3468	S1M10000046E01 S1M10000046E02	Staphylococcus aureus
3469	S1M10000046E04	Staphylococcus aureus
3470	S1M10000046E04	Staphylococcus aureus Staphylococcus aureus
3471	S1M1000046E07	Staphylococcus aureus Staphylococcus aureus
3472	S1M10000046E08	Staphylococcus aureus Staphylococcus aureus
3474	S1M1000046E10	Staphylococcus aureus
3475	S1M10000046F02	Staphylococcus aureus Staphylococcus aureus
3476	S1M1000046F05	Staphylococcus aureus
3477	S1M10000046F06	Staphylococcus aureus
3478	S1M10000046F08	Staphylococcus aureus
3479	S1M10000046F09	Staphylococcus aureus
3480	S1M10000046F10	Staphylococcus aureus
3481	S1M10000046F12	Staphylococcus aureus
3482	S1M10000046G01	Staphylococcus aureus
3483	S1M1000046G02	Staphylococcus aureus
3484	S1M10000046G03	Staphylococcus aureus
3485	S1M10000046G04	Staphylococcus aureus

SeqID	Clone name	Organism
3486	S1M10000046G07	Staphylococcus aureus
3487	S1M10000046G09	Staphylococcus aureus
3488	S1M10000046G10	Staphylococcus aureus
3489	S1M10000046H01	Staphylococcus aureus
3490	S1M10000046H10	Staphylococcus aureus
3491	S1M10000047A03	Staphylococcus aureus
3492	S1M10000047A04	Staphylococcus aureus
3493	S1M10000047A05	Staphylococcus aureus
3494	S1M10000047A06	Staphylococcus aureus
3495	S1M10000047A07	Staphylococcus aureus
3496	S1M10000047A08	Staphylococcus aureus
3497	S1M10000047A09	Staphylococcus aureus
3498	S1M10000047A10	Staphylococcus aureus
3499	S1M10000047A11	Staphylococcus aureus
3500	S1M10000047A12	Staphylococcus aureus
3501	S1M10000047B02	Staphylococcus aureus
3502	S1M10000047B04	Staphylococcus aureus
3503	S1M10000047B05	Staphylococcus aureus
3504	S1M10000047B06	Staphylococcus aureus
3505	S1M10000047B08	Staphylococcus aureus
3506	\$1M10000047B09	Staphylococcus aureus
3507	S1M10000047B10	Staphylococcus aureus
3508	S1M10000047B12	Staphylococcus aureus
3509	S1M10000047C01	Staphylococcus aureus
3510	S1M10000047C02	Staphylococcus aureus
3511	S1M10000047C03	Staphylococcus aureus
3512	S1M10000047C04	Staphylococcus aureus
3513	S1M1000047C06	Staphylococcus aureus
3514 3515	S1M10000047C08	Staphylococcus aureus
3515	S1M10000047C09 S1M10000047C11	Staphylococcus aureus
3516	S1M10000047C11 S1M10000047C12	Staphylococcus aureus
3518	S1M10000047C12	Staphylococcus aureus
3519	S1M1000047D02	Staphylococcus aureus
3520	S1M1000047D03	Staphylococcus aureus Staphylococcus aureus
3521	S1M1000047D04	Staphylococcus aureus
3522	S1M1000047D09	Staphylococcus aureus
3523	S1M10000047D09	Staphylococcus aureus Staphylococcus aureus
3524	S1M10000047D10	Staphylococcus aureus Staphylococcus aureus
3525	S1M10000047D12	Staphylococcus aureus
3526	S1M1000047E01	Staphylococcus aureus
3527	S1M1000047E02	Staphylococcus aureus
3528	S1M10000047E03	Staphylococcus aureus
3529	S1M10000047E04	Staphylococcus aureus
3530	S1M10000047E05	Staphylococcus aureus
3531	S1M10000047E06	Staphylococcus aureus
3532	S1M10000047E08	Staphylococcus aureus
3533	S1M10000047E09	Staphylococcus aureus
3534	S1M10000047E10	Staphylococcus aureus
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SeqID	Clone name	Organism
3535	S1M10000047E11	Staphylococcus aureus
3536	S1M10000047E12	Staphylococcus aureus
3537	S1M10000047F02	Staphylococcus aureus
3538	S1M10000047F03	Staphylococcus aureus
3539	S1M10000047F04	Staphylococcus aureus
3540	S1M10000047F05	Staphylococcus aureus
3541	S1M10000047F06	Staphylococcus aureus
3542	S1M10000047F07	Staphylococcus aureus
3543	S1M10000047F08	Staphylococcus aureus
3544	S1M10000047F09	Staphylococcus aureus
3545	S1M10000047F10	Staphylococcus aureus
3546	S1M10000047F11	Staphylococcus aureus
3547	S1M10000047F12	Staphylococcus aureus
3548	S1M10000047G01	Staphylococcus aureus
3549	S1M10000047G02	Staphylococcus aureus
3550	S1M10000047G04	Staphylococcus aureus
3551	S1M10000047G05	Staphylococcus aureus
3552	\$1M10000047G06	Staphylococcus aureus
3553	S1M10000047G07	Staphylococcus aureus
3554	S1M10000047G08	Staphylococcus aureus
3555	S1M10000047G09	Staphylococcus aureus
3556	S1M10000047G10	Staphylococcus aureus
3557	S1M10000047H03	Staphylococcus aureus
3558	S1M10000047H04	Staphylococcus aureus
3559	S1M10000047H05	Staphylococcus aureus
3560	S1M10000047H06	Staphylococcus aureus
3561	S1M10000047H07	Staphylococcus aureus
3562	S1M10000047H08	Staphylococcus aureus
3563	S1M10000047H09	Staphylococcus aureus
3564	S1M10000047H11	Staphylococcus aureus
3565 3566	S1M10000048A02	Staphylococcus aureus
3567	S1M10000048A03	Staphylococcus aureus
3568	S1M10000048A04 S1M10000048A05	Staphylococcus aureus
3569	S1M10000048A05	Staphylococcus aureus
3570	S1M1000048A08	Staphylococcus aureus
3571	S1M10000048A07	Staphylococcus aureus
3572	S1M1000048A09	Staphylococcus aureus
3573	S1M10000048A10	Staphylococcus aureus Staphylococcus aureus
3574	S1M1000048A11	
3575	S1M1000048A12	Staphylococcus aureus Staphylococcus aureus
3576	S1M1000048B02	Staphylococcus aureus
3577	S1M1000048B08	Staphylococcus aureus Staphylococcus aureus
3578	S1M1000048B08	Staphylococcus aureus
3579	S1M10000048B10	Staphylococcus aureus Staphylococcus aureus
3580	S1M1000048B12	Staphylococcus aureus Staphylococcus aureus
3581	S1M10000048B12	Staphylococcus aureus
3582	S1M1000048C02	Staphylococcus aureus
3583	S1M1000048C02	Staphylococcus aureus
3303	0.1.110000040000	propriyed coccus aureus

SeqID	Clone name	Organism
3584	S1M10000048C05	Staphylococcus aureus
3585	S1M10000048C06	Staphylococcus aureus
3586	S1M1000048C07	Staphylococcus aureus
3587	S1M1000048C08	Staphylococcus aureus
3588	S1M1000048C09	Staphylococcus aureus
3589	S1M10000048C11	Staphylococcus aureus
3590	S1M10000048D02	Staphylococcus aureus
3591	S1M10000048D08	Staphylococcus aureus
3592	S1M1000048D09	Staphylococcus aureus
3593	S1M10000048D10	Staphylococcus aureus
3594	S1M1000048D12	Staphylococcus aureus
3595	S1M1000048E02	Staphylococcus aureus
3596	S1M10000048E03	Staphylococcus aureus
3597	S1M10000048E04	Staphylococcus aureus
3598	S1M1000048E06	Staphylococcus aureus
3599	S1M10000048E07	Staphylococcus aureus
3600	S1M1000048E08	Staphylococcus aureus
3601	S1M1000048E10	Staphylococcus aureus
3602	S1M1000048E10	Staphylococcus aureus
3603	S1M10000048F07	Staphylococcus aureus
3604	S1M1000048F07	Staphylococcus aureus Staphylococcus aureus
3605	S1M1000048F09	Staphylococcus aureus Staphylococcus aureus
3606	S1M10000048F11	Staphylococcus aureus Staphylococcus aureus
3607	S1M10000048F12	Staphylococcus aureus
3608	S1M1000048G02	Staphylococcus aureus Staphylococcus aureus
3609	\$1M1000048G03	Staphylococcus aureus
3610	S1M1000048G04	Staphylococcus aureus
3611	S1M1000048G05	Staphylococcus aureus
3612	S1M1000048G07	Staphylococcus aureus
3613	S1M1000048G10	Staphylococcus aureus
3614	S1M1000048G11	Staphylococcus aureus
3615	S1M10000048H01	Staphylococcus aureus
3616	S1M10000048H02	Staphylococcus aureus
3617	S1M10000048H03	Staphylococcus aureus
3618	S1M10000048H04	Staphylococcus aureus
3619	S1M10000048H05	Staphylococcus aureus
3620	S1M10000048H07	Staphylococcus aureus
3621	S1M10000048H08	Staphylococcus aureus
3622	S1M10000048H09	Staphylococcus aureus
3623	S1M10000048H10	Staphylococcus aureus
3624	S1M10000048H11	Staphylococcus aureus
3625	S1M1000009E10	Staphylococcus aureus
3626	S1M10000001F01	Staphylococcus aureus
3627	S1M10000001101	Staphylococcus aureus
3628	S1M1000000B12 S1M10000003D09	Staphylococcus aureus Staphylococcus aureus
3629	S1M10000003D09	Staphylococcus aureus Staphylococcus aureus
3630	S1M10000001D11	Staphylococcus aureus
3631	S1M10000003B07	Staphylococcus aureus
3632	S1M10000002A07	Staphylococcus aureus
3032	DIMITO000003111	Suprificoccus aureus

SeqID	Clone name	Organism
3633	S1M10000047C07	Staphylococcus aureus
3634	S1M10000013F10	Staphylococcus aureus
3635	S1M10000014D11	Staphylococcus aureus
3636	S1M10000015F05	Staphylococcus aureus
3637	S1M10000048D01	Staphylococcus aureus
3638	S1M10000011C03	Staphylococcus aureus
3639	S1M10000012F03	Staphylococcus aureus
3640	S1M10000002F07	Staphylococcus aureus
3641	S1M10000048G01	Staphylococcus aureus
3642	S1M10000009G12	Staphylococcus aureus
3643	S1M10000012D05	Staphylococcus aureus
3644	S1M10000014D07	Staphylococcus aureus
3645	S1M10000047C05	Staphylococcus aureus
3646	S1M10000018D08*	Staphylococcus aureus
3647	S1M10000047B01	Staphylococcus aureus
3648	S1M10000047H10	Staphylococcus aureus
3649	S1M10000001A04	Staphylococcus aureus
3650	S1M10000016E01	Staphylococcus aureus
3651	S1M10000017E12	Staphylococcus aureus
3652	S1M10000019B01	Staphylococcus aureus
3653	S1M10000048F03	Staphylococcus aureus
3654	S1M10000034A07	Staphylococcus aureus
3655	S1M10000023G01	Staphylococcus aureus
3656	S1M10000021G12	Staphylococcus aureus
3657	S1M10000024E04	Staphylococcus aureus
3658	S1M10000028H08	Staphylococcus aureus
3659	S1M10000022B07	Staphylococcus aureus
3660 3661	S1M10000003A05	Staphylococcus aureus
3662	S1M10000003A09 S1M10000003E01	Staphylococcus aureus
3663	S1M10000003E01	Staphylococcus aureus
3664	S1M10000004C11	Staphylococcus aureus
3665	S1M1000007E08	Staphylococcus aureus
3666	S1M1000021G00	Staphylococcus aureus Staphylococcus aureus
3667	S1M10000024C00	Staphylococcus aureus
3668	S1M1000024D01	Staphylococcus aureus
3669	S1M1000027E03	Staphylococcus aureus
3670	S1M10000027G01	Staphylococcus aureus
3671	S1M1000029A03	Staphylococcus aureus
3672	S1M10000032B10	Staphylococcus aureus
3673	S1M10000032C07	Staphylococcus aureus
3674	S1M10000038D04	Staphylococcus aureus
3675	S1M10000047D07	Staphylococcus aureus
3676	S1M10000048B03	Staphylococcus aureus
3677	S1M10000048B06	Staphylococcus aureus
3678	S1M10000048C10	Staphylococcus aureus
3679	S1M10000048F05	Staphylococcus aureus
3680	S4M10000001C01	Salmonella typhimurium
3681	S4M10000002B06	Salmonella typhimurium

SeqID	Clone name	Organism
3682	S4M10000002B09	Salmonella typhimurium
3683	S4M1000002G04	Salmonella typhimurium
3684	S4M10000002G08	Salmonella typhimurium
3685	S4M1000005G05	Salmonella typhimurium
3686	S4M10000005H02	Salmonella typhimurium
3687	S4M1000006A06	Salmonella typhimurium
3688	S4M1000006A08	Salmonella typhimurium
3689	S4M10000006C05	Salmonella typhimurium
3690	S4M10000006F08	Salmonella typhimurium
3691	S4M10000007G01	Salmonella typhimurium
3692	S4M10000008C08	Salmonella typhimurium
3693	S4M10000008H10	Salmonella typhimurium
3694	S4M10000009A05	Salmonella typhimurium
3695	S4M10000010B05	Salmonella typhimurium
3696	S4M10000010D04	Salmonella typhimurium
3697	S4M10000010H04	Salmonella typhimurium
3698	S4M10000011D08	Salmonella typhimurium
3699	S4M10000011E08	Salmonella typhimurium
3700	S4M10000012B06	Salmonella typhimurium
3701	S4M10000012B12	Salmonella typhimurium
3702	S4M10000012D02	Salmonella typhimurium
3703	S4M10000013H02	Salmonella typhimurium
3704	S4M10000014B05	Salmonella typhimurium
3705	S4M10000014D04	Salmonella typhimurium
3706	S4M10000014D07	Salmonella typhimurium
3707	S4M10000014H02	Salmonella typhimurium
3708	S4M10000015B11	Salmonella typhimurium
3709	S4M10000015E09	Salmonella typhimurium
3710 3711	S4M10000016A02 S4M10000018D09	Salmonella typhimurium
3712	S4M10000018D09	Salmonella typhimurium
3712	S4M10000018E10	Salmonella typhimurium Salmonella typhimurium
3714	S4M10000018F10	
3715	S4M10000018G03	Salmonella typhimurium Salmonella typhimurium
3716	S4M10000019F05	Salmonella typhimurium
3717	S4M10000019F03	Salmonella typhimurium
3718	S4M1000019G05	Salmonella typhimurium
3719	S4M10000019H06	Salmonella typhimurium
3720	S4M10000020A04	Salmonella typhimurium
3721	S4M10000020F05	Salmonella typhimurium
3722	S4M10000020G10	Salmonella typhimurium
3723	S4M10000022D04	Salmonella typhimurium
3724	S4M10000022D12	Salmonella typhimurium
3725	S4M10000022E12	Salmonella typhimurium
3726	S4M10000022G07	Salmonella typhimurium
3727	S4M10000022H06	Salmonella typhimurium
3728	S4M10000023F01	Salmonella typhimurium
3729	S4M10000024B02	Salmonella typhimurium
3730	S4M10000024C06	Salmonella typhimurium
		<u> </u>

WO 01/70955 <u>TABLE IA</u> PCT/US01/09180

SeqID	Clone name	Organism
3731	S4M10000024C11	Salmonella typhimurium
3732	S4M10000024F08	Salmonella typhimurium
3733	S4M10000024G01	Salmonella typhimurium
3734	S4M10000024G04	Salmonella typhimurium
3735	S4M10000024G09	Salmonella typhimurium
3736	S4M10000024H02	Salmonella typhimurium
3737	S4M10000025A11	Salmonella typhimurium
3738	S4M10000025E02	Salmonella typhimurium
3739	S4M10000025E05	Salmonella typhimurium
3740	S4M10000025H07	Salmonella typhimurium
3741	S4M10000026C10	Salmonella typhimurium
3742	S4M10000026D04	Salmonella typhimurium
3743	S4M10000026E06	Salmonella typhimurium
3744	S4M10000026E12	Salmonella typhimurium
3745	S4M10000027C10	Salmonella typhimurium
3746	S4M10000027E02	Salmonella typhimurium
3747	S4M10000029B12	Salmonella typhimurium
3748	S4M10000029D12	Salmonella typhimurium
3749	S4M10000030D03	Salmonella typhimurium
3750	S4M10000030F07	Salmonella typhimurium
3751	S4M10000030G11	Salmonella typhimurium
3752	S4M10000032B12	Salmonella typhimurium
3753	S4M10000033F08	Salmonella typhimurium
3754	S4M10000033G05	Salmonella typhimurium
3755	S4M1000033G09	Salmonella typhimurium
3756	S4M1000034A02	Salmonella typhimurium
3757 3758	S4M1000034A09	Salmonella typhimurium
3759	S4M10000034D06 S4M10000034H05	Salmonella typhimurium Salmonella typhimurium
3760	S4M10000034H05 S4M10000034H09	Salmonella typnimurium Salmonella typhimurium
3761	S4M10000034H09 S4M10000035B01	Salmonetta typnimurtum Salmonella typhimurium
3762	S4M1000033B01	Salmonella typhimurium Salmonella typhimurium
3763	S4M1000033D01	Salmonella typhimurium Salmonella typhimurium
3764	S4M1000035D02	Salmonella typhimurium
3765	S4M1000033E03	Salmonella typhimurium
3766	S4M10000035F09	Salmonella typhimurium
3767	S4M1000035107	Salmonella typhimurium
3768	S4M1000036F07	Salmonella typhimurium
3769	S4M10000037A04	Salmonella typhimurium
3770	S4M10000037A10	Salmonella typhimurium
3771	S4M10000037E10	Salmonella typhimurium
3772	S4M10000037H09	Salmonella typhimurium
3773	S4M10000001H01	Salmonella typhimurium
3774	S4M10000002F06	Salmonella typhimurium
3775	S4M1000008D01	Salmonella typhimurium
3776	S4M10000009G11	Salmonella typhimurium
3777	S4M10000011F09	Salmonella typhimurium
3778	S4M10000020F08	Salmonella typhimurium
3779	S4M10000021E07	Salmonella typhimurium

SeqID	Clone name	Organism
3780	S4M10000022B05	Salmonella typhimurium
3781	S4M10000025H11	Salmonella typhimurium
3782	S4M10000026B10	Salmonella typhimurium
3783	S4M10000026E03	Salmonella typhimurium
3784	S4M10000029A03	Salmonella typhimurium
3785	S4M10000029C11	Salmonella typhimurium
3786	S4M10000030F06	Salmonella typhimurium
3787	S4M10000032F03	Salmonella typhimurium
3788	S4M10000032G01	Salmonella typhimurium
3789	S4M10000034C05	Salmonella typhimurium
3790	S4M10000034H04	Salmonella typhimurium
3791	S4M10000035A09	Salmonella typhimurium
3792	S4M10000035B06	Salmonella typhimurium
3793	S4M10000035F01	Salmonella typhimurium
3794	S4M10000037A08	Salmonella typhimurium
3795	S4M10000037E03	Salmonella typhimurium

TABLE IA

TABLE IB

Clone name	Clone SegID	PathoSeq Locus	Gene SegID (protein)	Genemarked gene	full length ORF Protein Seq
E3M10000001A02	8	EFA101409	4934	EFA1c0022 orf 11p	10524
E3M10000001A06	9	EFA100642	4884	EFA1c0041 orf 56p	10792
E3M10000001B01 .	10	EFA101409	4934	EFA1c0022 orf 11p	10524
E3M10000001B02	11	EFA100739	4888	EFA1c0022 orf 23p	10537
E3M10000001B02	11	EFA102549	5000	EFA1c0022 orf 24p	10538
E3M10000001B02	11	EFA102551	5001	EFA1c0022 orf 25p	10539
E3M10000001B05	12	EFA101165	4922	EFA1c0022 orf 8p	10559
E3M10000001B06	13	EFA101164	4921	EFA1c0022_orf_7p	10558
E3M10000001B08	14	EFA100642	4884	EFA1c0041_orf_56p	10792
E3M10000001B10	15	EFA101409	4934	EFA1c0022_orf_11p	10524
E3M10000001C02	16	EFA103038	5017	EFA1c0030_orf_17p	10613
E3M10000001C09	17	EFA103021	5015	EFA1c0030_orf_16p	10612
E3M10000001D02	18	EFA101159	4916	EFA1c0022_orf_2p	10543
E3M10000001D04	19	EFA100742	4891	EFA1c0022_orf_20p	10534
E3M10000001D04	19	EFA101417	4942	EFA1c0022_orf_18p	10531
E3M10000001D04	19	EFA102554	5002	EFA1c0022_orf_19p	10532
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E3M10000001D09	21	EFA100211	4871	EFA1c0022_orf_10p	10523
E3M10000001E01	22	EFA101162	4919	EFA1c0022_orf_5p	10555
E3M10000001E01	22	EFA101163	4920	EFA1c0022_orf_6p	10557
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E3M10000001E03	24	EFA100211	4871	EFA1c0022_orf_10p	10523
E3M10000001E04	25	EFA100642	4884	EFA1c0041_orf_56p	10792
E3M10000001E08	26	EFA102502	4995	EFA1c0031_orf_36p	10627
E3M10000001E09	27	EFA100210	4870	EFA1c0022_orf_9p	10560
E3M10000001E09	27	EFA100211	4871	EFA1c0022_orf_10p	10523
E3M10000001F02	28	EFA102502	4995	EFA1c0031_orf_36p	10627
E3M10000001F04	29	EFA102541	4998	EFA1c0028_orf_3p	10602
E3M10000001F06	30	EFA100642	4884	EFA1c0041_orf_56p	10792
E3M10000001F07	31	EFA101164	4921	EFA1c0022_orf_7p	10558
E3M10000001G02	32	EFA101409	4934	EFA1c0022_orf_11p	10524
E3M10000001G03	33	EFA100210	4870	EFA1c0022_orf_9p	10560
E3M10000001G03	33	EFA100211	4871	EFA1c0022_orf_10p	10523
E3M10000001G04	34	EFA101165	4922	EFA1c0022_orf_8p	10559
E3M10000001G05	35	EFA101160	4917	EFA1c0022_orf_3p	10549
E3M10000001H02	36	EFA102541	4998	EFA1c0028_orf_3p	10602
E3M10000001H03	37	EFA100210	4870	EFA1c0022_orf_9p	10560
E3M10000001H03	37	EFA100211	4871	EFA1c0022_orf_10p	10523
E3M10000001H04	38	EFA100742	4891	EFA1c0022_orf_20p	10534
E3M10000001H04	38	EFA101417	4942	EFA1c0022_orf_18p	10531
E3M10000001H04	38	EFA102554	5002	EFA1c0022_orf_19p	10532
E3M10000004A04	39	EFA101417	4942	EFA1c0022_orf_18p	10531
E3M10000004A04	39	EFA102554	5002	EFA1c0022_orf_19p	10532

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
E3M10000004C03	40	EFA100478	4880	EFA1c0012_orf_2p	10486
E3M10000004D01	41,	EFA101412	4937	EFA1c0022_orf_14p	10527
E3M10000004D01	41	EFA101413	4938	#N/A	#N/A
E3M10000004D01	41	EFA101414	4939	EFA1c0022_orf_15p	10528
E3M10000004D02	42	EFA102022	4974	EFA1c0044_orf_106p	10881
E3M10000004D02	42	EFA102023	4975	EFA1c0044_orf_107p	10882
E3M10000004D10	43	EFA101162	4919	EFA1c0022_orf_5p	10555
E3M10000004D10	43	EFA101163	4920	EFA1c0022_orf_6p	10557
E3M10000004E11	44	EFA101086	4910	EFA1c0040_orf_90p	10763
E3M10000004F08	45	EFA102549	5000	EFA1c0022_orf_24p	10538
E3M10000004F08	45	EFA102551	5001	EFA1c0022_orf_25p	10539
E3M10000004F10	46	EFA101086	4910	EFA1c0040 orf 90p	10763
E3M10000004G01	47	EFA103021	5015	EFA1c0030 orf 16p	10612
E3M10000004H11	48	EFA102549	5000	EFA1c0022 orf 24p	10538
E3M10000004H11	48	EFA102551	5001	EFA1c0022 orf 25p	10539
E3M10000005A07	49	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000005B01	50	EFA101414	4939	EFA1c0022 orf 15p	10528
E3M10000005B01	50	EFA101415	4940	EFA1c0022 orf 16p	10529
E3M10000005B08	51	EFA102549	5000	EFA1c0022 orf 24p	10538
E3M10000005B08	51	EFA102551	5001	EFA1c0022 orf 25p	10539
E3M10000005C01	52	EFA103021	5015	EFA1c0030 orf 16p	10612
E3M10000005C03	53	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000005C04	54	EFA102186	4981	EFA1c0045 orf 94p	10949
E3M1000005C04	54	EFA102453	4993	EFA1c0045 orf 203p	10931
E3M1000005C04	54	EFA102728	5006	EFA1c0045_orf_93p	10948
E3M10000005D03	55	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000005D04	56	EFA103021	5015	EFA1c0030 orf 16p	10612
E3M10000005D10	57	EFA102549	5000	EFA1c0022 orf 24p	10538
E3M10000005D10	57	EFA102551	5001	EFA1c0022 orf 25p	10539
E3M10000005E01	58	EFA102549	5000	EFA1c0022 orf 24p	10538
E3M10000005E01	58	EFA102551	5001	EFA1c0022 orf 25p	10539
E3M10000005E02	59	EFA102549	5000	EFA1c0022 orf 24p	10538
E3M10000005E02	59	EFA102551	5001	EFA1c0022 orf 25p	10539
E3M10000005E03	60	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000005E08	61	EFA101403	4932	EFA1c0033 orf 54p	10662
E3M10000005F07	62	EFA103021	5015	EFA1c0030 orf 16p	10612
E3M10000005F10	63	EFA102549	5000	EFA1c0022 orf 24p	10538
E3M10000005F10	63	EFA102551	5001	EFA1c0022 orf 25p	10539
E3M10000005G05	64	EFA102549	5000	EFA1c0022 orf 24p	10538
E3M10000005G05	64	EFA102551	5001	EFA1c0022_orf_25p	10539
E3M10000005H04	65	EFA103021	5015	EFA1c0030_orf_16p	10612
E3M10000005H04	66	EFA101162	4919	EFA1c0022 orf 5p	10512
E3M10000006B03	66	EFA101163	4920	EFA1c0022_orf 6p	10557
E3M10000006C01	67	EFA101416	4941	EFA1c0022_orf_17p	10537
E3M10000006C01	67	EFA101417	4942	EFA1c0022_orf_17p	10530
E3M10000006C01	68	EFA102549	5000	EFA1c0022_orf_18p	10531
E3M10000006C12	68	EFA102551	5001	EFA1c0022_orf_24p	10538
E3M10000006C12	69	EFA101416	4941	EFA1c0022_orf_25p	10539

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
E3M10000006D03	69	EFA101417	4942	EFA1c0022_orf_18p	10531
E3M10000006E11	70	EFA102541	4998	EFA1c0028_orf_3p	10602
E3M10000006E11	70	EFA102542	4999	EFA1c0028_orf_4p	10603
E3M10000006F04	71	EFA102541	4998	EFA1c0028_orf_3p	10602
E3M10000006F04	71	EFA102542	4999	EFA1c0028_orf_4p	10603
E3M10000006G04	72	EFA101162	4919	EFA1c0022_orf_5p	10555
E3M10000006G04	72	EFA101163	4920	EFA1c0022_orf_6p	10557
E3M10000006G12	73	EFA101162	4919	EFA1c0022_orf_5p	10555
E3M10000006G12	73	EFA101163	4920	EFA1c0022_orf_6p	10557
E3M10000006H09	74	EFA100642	4884	EFA1c0041_orf_56p	10792
E3M10000007A02	75	EFA101162	4919	EFA1c0022_orf_5p	10555
E3M10000007A02	75	EFA101163	4920	EFA1c0022 orf 6p	10557
E3M10000007B02	76	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000007B02	76	EFA101163	4920	EFA1c0022 orf 6p	10557
E3M10000007B03	77	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000007B03	77	EFA101163	4920	EFA1c0022 orf 6p	10557
E3M10000007C03	78	EFA101416	4941	EFA1c0022 orf 17p	10530
E3M10000007C03	78	EFA101417	4942	EFA1c0022 orf 18p	10531
E3M10000007C04	79	EFA100642	4884	EFA1c0041 orf 56p	10792
E3M10000007D03	80	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000007D03	80	EFA101163	4920	EFA1c0022 orf 6p	10557
E3M10000007E05	81	EFA100742	4891	EFA1c0022 orf 20p	10534
E3M10000007E05	81	EFA101417	4942	EFA1c0022 orf 18p	10531
E3M10000007E05	81	EFA102554	5002	EFA1c0022 orf 19p	10532
E3M10000007F01	82	EFA101162	4919	EFA1c0022_orf_5p	10555
E3M10000007F01	82	EFA101163	4920	EFA1c0022 orf 6p	10557
E3M10000007F06	83	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000007F06	83	EFA101163	4920	EFA1c0022 orf 6p	10557
E3M1000007G01	84	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000007G01	84	EFA101163	4920	EFA1c0022 orf 6p	10557
E3M10000008C03	85	EFA102501	4994	EFA1c0031 orf 35p	10626
E3M10000008C08	86	EFA101536	4946	EFA1c0042 orf 46p	10823
E3M10000008C09	87	EFA101410	4935	EFA1c0022 orf 12p	10525
E3M10000008D08	88	EFA102501	4994	EFA1c0031 orf 35p	10626
E3M10000008E02	89	EFA100783	4895	EFA1c0042 orf 141p	10811
E3M10000008G05	90	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000008G05	90	EFA101163	4920	EFA1c0022 orf 6p	10557
E3M10000008G09	91	EFA103021	5015	EFA1c0030 orf 16p	10612
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E3M10000008H02	92	EFA101695	4954	EFA1c0030_on_17p	10629
E3M10000000102	93	EFA103508	5029	EFA1c0033 orf 95p	10672
E3M10000009C09	94	EFA100870	4899	EFA1c0031_orf_36p	10627
E3M10000009C09	95	EFA101410	4935	EFA1c0031_orf_12p	10527
E3M10000009E02	96	EFA101410	4935	EFA1c0022_orf_12p	10525
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	98	EFA101160 EFA102501	4917	EFA1c0022_ort_3p EFA1c0031 orf 35p	10549
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E3M10000010D05	101	EFA100757	4894	EFA1c0044_orf_27p	10897
E3M10000010F01	102	EFA102551	5001	EFA1c0022_orf_25p	10539
E3M10000010G05	103	EFA101164	4921	EFA1c0022_orf_7p	10558
E3M10000010G07	104	EFA101165	4922	EFA1c0022_orf_8p	10559
E3M10000010G09	105	EFA103571	5030	EFA1c0044_orf_101p	10879
E3M10000010G10	106	EFA102091	4977	EFA1c0010_orf_3p	10481
E3M10000010H02	107	EFA100194	4868	EFA1c0022_orf_26p	10540
E3M10000011A09	108	EFA103038	5017	EFA1c0030_orf_17p	10613
E3M10000011B03	109	EFA102091	4977	EFA1c0010_orf_3p	10481
E3M10000011B09	110	EFA102501	4994	EFA1c0031_orf_35p	10626
E3M10000011C07	111	EFA101790	4959	EFA1c0042 orf 111p	10803
E3M10000011D03	112	EFA100210	4870	EFA1c0022 orf 9p	10560
E3M10000011D03	112	EFA100211	4871	EFA1c0022 orf 10p	10523
E3M10000011H02	113	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000011H05	114	EFA101164	4921	EFA1c0022 orf 7p	10558
E3M10000012B01	115	EFA100642	4884	EFA1c0041 orf 56p	10792
E3M10000012B02	116	EFA100151	4864	EFA1c0021 orf 14p	10516
E3M10000012B07	117	EFA101410	4935	EFA1c0022 orf 12p	10525
E3M10000012B07	117	EFA101411	4936	EFA1c0022 orf 13p	10526
E3M10000012B07	117	EFA101412	4937	EFA1c0022 orf 14p	10527
E3M10000012B08	118	EFA101409	4934	EFA1c0022 orf 11p	10524
E3M10000012C01	119	EFA100642	4884	EFA1c0041 orf 56p	10792
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E3M10000012F05	122	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000012F06	123	EFA101409	4934	EFA1c0022 orf 11p	10524
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E3M10000013A06	128	EFA101159	4916	EFA1c0022 orf 2p	10543
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E3M10000013D02	131	EFA101160	4917	EFA1c0022 orf 3p	10549
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E3M10000013D10	133	EFA100210	4870	EFA1c0022_orf_9p	10560
E3M10000013D10	133	EFA100211	4871	EFA1c0022_orf_10p	10523
E3M10000013E02	134	EFA100642	4884	EFA1c0041 orf 56p	10792
E3M10000013E08	135	EFA102501	4994	EFA1c0031 orf 35p	10626
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E3M10000013F12	137	EFA101164	4921	EFA1c0022 orf 7p	10558
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E3M10000013H10	141	EFA101164	4921	EFA1c0022 orf 7p	10558
E3M10000014B12	142	EFA100739	4888	EFA1c0022 orf 23p	10537
E3M10000014B12	142	EFA102549	5000	EFA1c0022 orf 24p	10538
E3M10000014B12	142	EFA102551	5001	EFA1c0022_orf_25p	10539
E3M10000014E12	143	EFA101409	4934	EFA1c0022 orf 11p	10524
E3M10000014E12	143	EFA101410	4935	EFA1c0022 orf 12p	10525
E3M10000014G09	144	EFA100991	4905	EFA1c0035 orf 60p	10681
E3M10000014G09	144	EFA103033	5016	EFA1c0035 orf 60p	10681
E3M10000015B04	145	EFA100065	4863	EFA1c0042_orf_14p	10813
E3M10000015B12	146	EFA101162	4919	EFA1c0022_orf_5p	10515
E3M10000015E12	147	EFA100210	4870	EFA1c0022_orf_9p	10560
E3M10000015E12	147	EFA100211	4871	EFA1c0022_orf_10p	10523
E3M10000015E12	148	EFA101753	4957	EFA1c0022_orf_10p	10523
E3M10000016A04	149	EFA101409	4934	EFA1c0022_orf_11p	10524
E3M10000016A04	150	EFA101163	4920	EFA1c0022_orf_fp	10524
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E3M10000016C11	151	EFA101164 EFA102774	5009	EFA1c0022_0ff_/p	10338
E3M10000016E03	152	EFA102774 EFA102205	4983	EFA1c0044_orf_25p	
E3M10000016F06	153	EFA101410	4983		10769
	153			EFA1c0022_orf_12p	10525
E3M10000016F10 E3M10000016H05		EFA101411	4936	EFA1c0022_orf_13p	10526
	154 155	EFA101160	4917	EFA1c0022_orf_3p	10549
E3M10000016H10 E3M10000017A09		EFA101409	4934	EFA1c0022_orf_11p	10524
E3M10000017A09	156 156	EFA101161	4918	EFA1c0022_orf_4p	10551
E3M10000017A09		EFA101162	4919 4937	EFA1c0022_orf_5p	10555
E3M10000017D09	157 158	EFA101412 EFA102091		EFA1c0022_orf_14p	10527
			4977	EFA1c0010_orf_3p	10481
E3M10000018C02	159	EFA100642	4884	EFA1c0041_orf_56p	10792
E3M10000018E01	160	EFA103021	5015	EFA1c0030_orf_16p	10612
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E3M10000018H06	162	EFA101417	4942	EFA1c0022_orf_18p	10531
E3M10000019B06	163	EFA100151	4864	EFA1c0021_orf_14p	10516
E3M10000019D02	164	EFA102022	4974	EFA1c0044_orf_106p	10881
E3M10000019E03	165	EFA100870	4899	EFA1c0031_orf_36p	10627
E3M10000019E03	165	EFA102502	4995	EFA1c0031_orf_36p	10627
E3M10000019E04	166	EFA102501	4994	EFA1c0031_orf_35p	10626
E3M10000020G04	167	EFA100870	4899	EFA1c0031_orf_36p	10627
E3M10000020G04	167	EFA102502	4995	EFA1c0031_orf_36p	10627
E3M10000020H05	168	EFA103038	5017	EFA1c0030_orf_17p	10613
E3M10000021A08	169	EFA101162	4919	EFA1c0022_orf_5p	10555
E3M10000021A08	169	EFA101163	4920	EFA1c0022_orf_6p	10557
E3M10000021A11	170	EFA101417	4942	EFA1c0022_orf_18p	· 10531
E3M10000021B10	171	EFA101163	4920	EFA1c0022_orf_6p	10557
E3M10000021C03	172	EFA102501	4994	EFA1c0031_orf_35p	10626
E3M10000021C04	173	EFA101161	4918	EFA1c0022_orf_4p	10551
E3M10000021C08	174	EFA101160	4917	EFA1c0022_orf_3p	10549
E3M10000021D04	175	EFA100870	4899	EFA1c0031_orf_36p	10627

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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E3M10000021G10	178	EFA100642	4884	EFA1c0041 orf 56p	10792
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E3M10000021H11	180	EFA101163	4920	EFA1c0022_orf_6p	10557
E3M10000022A04	181	EFA101410	4935	EFA1c0022_orf_12p	10525
E3M10000022A11	182	EFA101417	4942	EFA1c0022_orf_18p	10531
E3M10000022B04	183	EFA101410	4935	EFA1c0022_orf_12p	10525
E3M10000022B05	184	EFA101410	4935	EFA1c0022 orf 12p	10525
E3M10000022B05	184	EFA101411	4936	EFA1c0022 orf 13p	10526
E3M10000022B07	185	EFA103571	5030	EFA1c0044_orf_101p	10879
E3M10000022C05	186	EFA101160	4917	EFA1c0022 orf 3p	10549
E3M10000022C05	186	EFA101161	4918	EFA1c0022 orf 4p	10551
E3M10000022C06	187	EFA100978	4904	EFA1c0022 orf 27p	10541
E3M10000022C09	188	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000022D04	189	EFA101412	4937	EFA1c0022_orf_14p	10527
E3M10000022F05	190	EFA102501	4994	EFA1c0031_orf_35p	10626
E3M10000022F06	191	EFA101161	4918	EFA1c0022 orf 4p	10551
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E3M10000023A03	195	EFA101413	4938	#N/A	#N/A
E3M10000023A06	196	EFA100978	4904	EFA1c0022 orf 27p	10541
E3M10000023A07	197	EFA102502	4995	EFA1c0031 orf 36p	10627
E3M10000023A09	198	EFA100704	4887	EFA1c0010 orf 4p	10482
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E3M10000023B02	199	EFA101160	4917	EFA1c0022 orf 3p	10549
E3M10000023B06	200	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000023C03	201	EFA101409	4934	EFA1c0022_orf_11p	10524
E3M10000023C03	201	EFA101410	4935	EFA1c0022 orf 12p	10525
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E3M10000023C08	204	EFA100955	4902	EFA1c0022 orf 28p	10542
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E3M10000023C09	205	EFA101160	4917	EFA1c0022_orf_3p	10549
E3M10000023D02	206	EFA102501	4994	EFA1c0031 orf 35p	10626
E3M10000023D04	207	EFA101160	4917	EFA1c0022 orf 3p	10549
E3M10000023D10	208	EFA101413	4938	#N/A	#N/A
E3M10000023E04	209	EFA101412	4937	EFA1c0022 orf 14p	10527
E3M10000023E07	210	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000023E09	211	EFA102501	4994	EFA1c0031_orf_35p	10626
E3M10000023F02	212	EFA101412	4937	EFA1c0022 orf 14p	10527
E3M10000023F10	213	EFA102551	5001	EFA1c0022_orf_25p	10539
E3M10000023T10	214	EFA101160	4917	EFA1c0022_orf 3p	10549
E3M10000023G02	215	EFA101414	4939	EFA1c0022_orf_15p	10528
E3M10000023G10	216	EFA101411	4936	EFA1c0022_orf_13p	10526
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E3M10000024A03	218	EFA101161	4918	EFA1c0022_orf_4p	10551
E3M10000024A04	219	EFA102006	4973	EFA1c0025_orf_17p	10580
E3M10000024A08	220	EFA101160	4917	EFA1c0022 orf 3p	10549
E3M10000024A08	220	EFA101161	4918	EFA1c0022 orf 4p	10551
E3M10000024C06	221	EFA102501	4994	EFA1c0031 orf 35p	10626
E3M10000025A06	222	EFA101160	4917	EFA1c0022 orf 3p	10549
E3M10000025B01	223	EFA100194	4868	EFA1c0022 orf 26p	10540
E3M10000025B01	223	EFA100978	4904	EFA1c0022 orf 27p	10541
E3M10000025B03	224	EFA101411	4936	EFA1c0022 orf 13p	10526
E3M10000025B03	224	EFA101412	4937	EFA1c0022 orf 14p	10527
E3M10000025B05	225	EFA100978	4904	EFA1c0022 orf 27p	10541
E3M10000025B10	226	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000025C01	227	EFA102502	4995	EFA1c0031 orf 36p	10627
E3M10000025C04	228	EFA101159	4916	EFA1c0022 orf 2p	10543
E3M10000025C05	229	EFA102549	5000	EFA1c0022_orf_24p	10538
E3M10000025C05	229	EFA102551	5001	EFA1c0022_orf_25p	10539
E3M10000025C07	230	EFA100642	4884	EFA1c0022_off_25p	10792
E3M10000025C07	231	EFA100870	4899	EFA1c0031 orf 36p	10/92
E3M10000025C08	231	EFA102502	4995		10627
E3M10000025C08	231	EFA102501	4993	EFA1:0031_orf_36p	10627
E3M10000025C11	232	EFA102301 EFA101162	4994	EFA1c0031_orf_35p	
E3M10000025C11	234			EFA1c0022_orf_5p	10555
E3M10000025D01	234	EFA101160 EFA101161	4917 4918	EFA1c0022_orf_3p	10549
				EFA1c0022_orf_4p	10551
E3M10000025D10	235	EFA102501	4994	EFA1c0031_orf_35p	10626
E3M10000025E07	236	EFA101165	4922	EFA1c0022_orf_8p	10559
E3M10000025E08	237	EFA100955	4902	EFA1c0022_orf_28p	10542
E3M10000025E12	238	EFA102728	5006	EFA1c0045_orf_93p	10948
E3M10000025F04	239	EFA101160	4917	EFA1c0022_orf_3p	10549
E3M10000025F04	239	EFA101161	4918	EFA1c0022_orf_4p	10551
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E3M10000025F08	241	EFA103038	5017	EFA1c0030_orf_17p	10613
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E3M10000025G02	246	EFA101164	4921	EFA1c0022_orf_7p	10558
E3M10000025G07	247	EFA101159	4916	EFA1c0022_orf_2p	10543
E3M10000025G09	248	EFA102185	4980	EFA1c0045_orf_95p	10950
E3M10000027A02	249	EFA101416	4941	EFA1c0022_orf_17p	10530
E3M10000027A07	250	EFA101160	4917	EFA1c0022_orf_3p	10549
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E3M10000027B07	252	EFA101163	4920	EFA1c0022 orf 6p	10557
E3M10000027B08	253	EFA101160	4917	EFA1c0022_orf 3p	10549
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E3M10000027D03	258	EFA102502	4995	EFA1c0031 orf 36p	10627
E3M10000027D05	259	EFA101162	4919	EFA1c0022 orf 5p	10555
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E3M10000027D10	261	EFA100704	4887	EFA1c0010 orf 4p	10482
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E3M10000028A02	266	EFA102554	5002	EFA1c0022 orf 19p	10532
E3M10000028A03	267	EFA102551	5001	EFA1c0022_orf_25p	10539
E3M10000028A04	268	EFA101410	4935	EFA1c0022 orf 12p	10525
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E3M10000028A08	271	EFA101424	4943	EFA1c0041 orf 39p	10784
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E3M10000028B02	273	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000028B02	273	EFA102542	4999	EFA1c0028 orf 4p	10603
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E3M10000028B05	276	EFA101424	4943	EFA1c0041 orf 39p	10784
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E3M10000028B06	277	EFA103038	5017	EFA1c0030 orf 17p	10613
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E3M10000028C02	281	EFA102541	4998	EFA1c0028 orf 3p	10602
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E3M10000028C04	282	EFA101322	4927	EFA1c0030 orf 57p	10620
E3M10000028C05	283	EFA101160	4917	EFA1c0022 orf 3p	10549
E3M10000028C06	284	EFA100151	4864	EFA1c0021 orf 14p	10516
E3M10000028C07	285	EFA101022	4906	EFA1c0043 orf 69p	10875
E3M10000028C07	286	EFA102541	4998	EFA1c0043_0ff_09p	10602
E3M10000028C08	286	EFA102542	4999	EFA1c0028_orf_4p	10603
E3M10000028C08	287	EFA100194	4868	EFA1c0028_off_4p	10540
E3M10000028D01	287	EFA100194 EFA100978	4904	EFA1c0022_orf_2op	10540
E3M10000028D01	288	EFA101022	4904	EFA1c0022_orf_2/p EFA1c0043_orf_69p	10341
E3M10000028D02	288	EFA101022 EFA101080	4909	#N/A	#N/A
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Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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E3M10000028E01	292	EFA101121	4912	EFA1c0036_orf_112p	10686
E3M10000028E04	293	EFA101370	4931	EFA1c0040_orf_103p	10738
E3M10000028E07	294	EFA102541	4998	EFA1c0028_orf_3p	10602
E3M10000028F02	295	EFA101161	4918	EFA1c0022_orf_4p	10551
E3M10000028F03	296	EFA100742	4891	EFA1c0022_orf_20p	10534
E3M10000028F03	296	EFA101417	4942	EFA1c0022_orf_18p	10531
E3M10000028F03	296	EFA102554	5002	EFA1c0022_orf_19p	10532
E3M10000028F04	297	EFA101163	4920	EFA1c0022_orf_6p	10557
E3M10000028F04	297	EFA101164	4921	EFA1c0022_orf_7p	10558
E3M10000028F05	298	EFA100704	4887	EFA1c0010 orf 4p	10482
E3M10000028F06	299	EFA101164	4921	EFA1c0022 orf 7p	10558
E3M10000028F07	300	EFA100642	4884	EFA1c0041_orf_56p	10792
E3M10000028G05	301	EFA100704	4887	EFA1c0010 orf 4p	10482
E3M10000028G06	302	EFA100748	4892	EFA1c0011 orf 10p	10483
E3M10000028G07	303	EFA101410	4935	EFA1c0022 orf 12p	10525
E3M10000028G07	303	EFA101411	4936	EFA1c0022 orf 13p	10526
E3M10000028H04	304	EFA101409	4934	EFA1c0022 orf 11p	10524
E3M10000028H07	305	EFA103062	5019	EFA1c0030 orf 19p	10615
E3M10000029A02	306	EFA101160	4917	EFA1c0022 orf 3p	10549
E3M10000029A04	307	EFA100210	4870	EFA1c0022 orf 9p	10560
E3M10000029A05	308	EFA100642	4884	EFA1c0041_orf_56p	10792
E3M10000029A10	309	EFA102091	4977	EFA1c0010 orf 3p	10481
E3M10000029A11	310	EFA101413	4938	#N/A	#N/A
E3M10000029B01	311	EFA103295	5024	EFA1c0032 orf 1p	10633
E3M10000029B02	312	EFA101160	4917	EFA1c0022 orf 3p	10549
E3M10000029B05	313	EFA103038	5017	EFA1c0030 orf 17p	10613
E3M10000029B06	314	EFA100914	4900	EFA1c0024 orf 9p	10579
E3M10000029B08	315	EFA102338	4987	EFA1c0032 orf 8p	10651
E3M10000029B11	316	EFA100397	4877	EFA1c0041 orf 148p	10773
E3M10000029B12	317	EFA100642	4884	EFA1c0041 orf 56p	10792
E3M10000029C01	318	EFA102541	4998	EFA1c0028_orf_3p	10602
E3M10000029C02	319	EFA102788	5011	EFA1c0033_orf_41p	10661
E3M10000029C03	320	EFA102253	4984	EFA1c0038 orf 85p	10727
E3M10000029C04	321	EFA102503	4996	EFA1c0032 orf 32p	10643
E3M10000029C05	322	EFA100399	4878	EFA1c0041 orf 104p	10766
E3M10000029C06	323	EFA101414	4939	EFA1c0022 orf 15p	10528
E3M10000029C06	323	EFA101415	4940	EFA1c0022 orf 16p	10529
E3M10000029C07	324	EFA102352	4990	EFA1c0032 orf 21p	10635
E3M10000029C07	324	EFA102353	4991	EFA1c0032 orf 22p	10636
E3M10000029C08	325	EFA101868	4966	EFA1c0042 orf 69p	10829
E3M10000029C09	326	EFA101121	4912	EFA1c0036 orf 112p	10686
E3M10000029C10	327	EFA102656	5004	EFA1c0039 orf 26p	10734
E3M10000029C12	328	EFA101121	4912	EFA1c0036 orf 112p	10686
E3M10000029C12	329	EFA101080	4909	#N/A	#N/A
E3M10000029D03	330	EFA101160	4917	EFA1c0022 orf 3p	10549
LIJ:VII 100000Z7IJUJ					
E3M10000029D04	331	EFA102656	5004	EFA1c0039 orf 26p	10734

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
E3M10000029D06	333	EFA100210	4870	EFA1c0022_orf_9p	10560
E3M10000029D06	333	EFA101165	4922	EFA1c0022_orf_8p	10559
E3M10000029D08	334	EFA102736	5007	EFA1c0022_orf_60p	10556
E3M10000029D12	335	EFA101410	4935	EFA1c0022 orf 12p	10525
E3M10000029E01	336	EFA101404	4933	EFA1c0033_orf_55p	10663
E3M10000029E02	337	EFA102051	4976	#N/A	#N/A
E3M10000029E03	338	EFA102502	4995	EFA1c0031 orf 36p	10627
E3M10000029E05	339	EFA101686	4953	EFA1c0045 orf 63p	10940
E3M10000029E07	340	EFA100919	4901	EFA1c0013 orf 12p	10491
E3M10000029E08	341	EFA101022	4906	EFA1c0043 orf_69p	10875
E3M10000029E09	342	EFA102656	5004	EFA1c0039 orf 26p	10734
E3M10000029E12	343	EFA100397	4877	EFA1c0041 orf 148p	10773
E3M10000029F01	344	EFA100023	4862	EFA1c0017 orf 1p	10505
E3M10000029F05	345	EFA102503	4996	EFA1c0032 orf 32p	10643
E3M10000029F06	346	EFA101795	4962	EFA1c0045 orf 165p	10922
E3M10000029F09	347	EFA100689	4886	EFA1c0038 orf 54p	10717
E3M10000029F10	348	EFA100919	4901	EFA1c0013 orf 12p	10491
E3M10000029F11	349	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000029F12	350	EFA102282	4985	EFA1c0038 orf 89p	10729
E3M10000029G01	351	EFA100394	4876	EFA1c0034 orf 6p	10675
E3M10000029G01	352	EFA102656	5004	EFA1c0034_orf_26p	10073
E3M10000029G04	353	EFA102351	4989	EFA1c0039_0ff_20p	10/34
E3M10000029G07	354	EFA101121	4912	EFA1c0036 orf 112p	10686
E3M10000029G07	355	EFA103571	5030	EFA1c0034 orf 101p	10879
E3M10000029G09	356	EFA102201	4982	#N/A	#N/A
E3M10000029G10	357	EFA101797	4963	EFA1c0045 orf 167p	10924
E3M10000029G10	358	EFA102006	4973	EFA1c0043_orf_16/p	10524
E3M10000029G11	359	EFA101541	4948	EFA1c0012 orf 5p	10380
E3M10000029G12	360	EFA101341 EFA101339			1
E3M10000029H02	360		4928	EFA1c0040_orf_13p	10743
		EFA101340	4929	EFA1c0040_orf_15p	10745
E3M10000029H04	361	EFA102352	4990	EFA1c0032_orf_21p	10635
E3M10000029H04	361	EFA102353	4991	EFA1c0032_orf_22p	10636
E3M10000029H05	362	EFA102091	4977	EFA1c0010_orf_3p	10481
E3M10000029H07	363	EFA100190	4867	EFA1c0010_orf_2p	10480
E3M10000029H08	364	EFA101416	4941	EFA1c0022_orf_17p	10530
E3M10000029H11	365	EFA101159	4916	EFA1c0022_orf_2p	10543
E3M10000030A05	366	EFA102501	4994	EFA1c0031_orf_35p	10626
E3M10000030A08	367	EFA102351	4989	EFA1c0032_orf_20p	10634
E3M10000030A09	368	EFA102501	4994	EFA1c0031_orf_35p	10626
E3M10000030A11	369	EFA102736	5007	EFA1c0022_orf_60p	10556
E3M10000030B03	370	EFA100704	4887	EFA1c0010_orf_4p	10482
E3M10000030B04	371	EFA103038	5017	EFA1c0030_orf_17p	10613
E3M10000030B05	372	EFA102656	5004	EFA1c0039_orf_26p	10734
E3M10000030B06	373	EFA101162	4919	EFA1c0022_orf_5p	10555
E3M10000030B07	374	EFA100642	4884	EFA1c0041_orf_56p	10792
E3M10000030B08	375	EFA102656	5004	EFA1c0039_orf_26p	10734
E3M10000030B10	376	EFA102655	5003	EFA1c0039_orf_25p	10733
E3M10000030B11	377	EFA101121	4912	EFA1c0036_orf_112p	10686

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
E3M10000030B12	378	EFA102352	4990	EFA1c0032_orf_21p	10635
E3M10000030B12	378	EFA102353	4991	EFA1c0032 orf 22p	10636
E3M10000030C03	379	EFA100151	4864	EFA1c0021 orf 14p	10516
E3M10000030C04	380	EFA101165	4922	EFA1c0022_orf_8p	10559
E3M10000030C12	381	EFA102351	4989	EFA1c0032 orf 20p	10634
E3M10000030D02	382	EFA102350	4988	EFA1c0032 orf 19p	10632
E3M10000030D05	383	EFA101414	4939	EFA1c0022_orf_15p	10528
E3M10000030D08	384	EFA102780	5010	EFA1c0045 orf 101p	10908
E3M10000030D09	385	EFA102780	5010	EFA1c0045 orf 101p	10908
E3M10000030D10	386	EFA102656	5004	EFA1c0039_orf_26p	10734
E3M10000030D12	387	EFA101417	4942	EFA1c0022 orf 18p	10531
E3M10000030E01	388	EFA101410	4935	EFA1c0022 orf 12p	10525
E3M10000030E01	388	EFA101411	4936	EFA1c0022 orf 13p	10526
E3M10000030E02	389	EFA100329	4875	EFA1c0041 orf 35p	10782
E3M10000030E04	390	EFA102655	5003	EFA1c0039 orf 25p	10733
E3M10000030E08	391	EFA101540	4947	EFA1c0012 orf 4p	10487
E3M10000030E09	392	EFA103365	5026	EFA1c0022 orf 1p	10533
E3M10000030E10	393	EFA102656	5004	EFA1c0039 orf 26p	10734
E3M10000030F01	394	EFA102655	5003	EFA1c0039_orf_25p	10733
E3M10000030F04	395	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000030F06	396	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000030F07	397	EFA102656	5004	EFA1c0039 orf 26p	10734
E3M10000030F10	398	EFA101417	4942	EFA1c0022 orf 18p	10531
E3M10000030F12	399	EFA102091	4977	EFA1c0010 orf 3p	10481
E3M10000030G01	400	EFA102551	5001	EFA1c0022 orf 25p	10539
E3M10000030G03	401	EFA100023	4862	EFA1c0017 orf 1p	10505
E3M10000030G06	402	EFA101686	4953	EFA1c0045 orf 63p	10940
E3M10000030G08	403	EFA102501	4994	EFA1c0031 orf 35p	10626
E3M10000030G09	404	EFA103210	5022	EFA1c0036 orf 119p	10688
E3M10000030G12	405	EFA103504	5028	EFA1c0033 orf_94p	10671
E3M10000030H03	406	EFA101258	4926	EFA1c0045 orf 160p	10918
E3M10000030H04	407	EFA101121	4912	EFA1c0036_orf_112p	10686
E3M10000030H06	408	EFA101161	4918	EFA1c0022 orf 4p	10551
E3M10000030H07	409	EFA101165	4922	EFA1c0022 orf 8p	10559
E3M10000030H08	410	EFA102501	4994	EFA1c0031 orf 35p	10626
E3M10000030H10	411	EFA102091	4977	EFA1c0010 orf 3p	10481
E3M10000030H11	412	EFA100615	4881	EFA1c0016 orf 29p	10501
E3M10000031A02	413	EFA102006	4973	EFAIc0025 orf 17p	10580
E3M10000031A06	414	EFA100970	4903	EFA1c0044 orf 98p	10906
E3M10000031A07	415	EFA102201	4982	#N/A	#N/A
E3M10000031A07	416	EFA100642	4884	EFA1c0041 orf 56p	10792
E3M10000031R08	417	EFA100289	4872	EFA1c0042 orf 139p	10810
E3M10000031B02	418	EFA100426	4879	EFA1c0036_orf_59p	10702
E3M10000031B03	419	EFA100394	4876	EFA1c0034 orf 6p	10675
E3M10000031B04	420	EFA102183	4979	EFA1c0045 orf 97p	10073
E3M10000031B10	421	EFA101253	4924	EFA1c0043_orf_178p	10852
E3M10000031B10	421	EFA100190	4867	EFA1c0010 orf 2p	10480
E3M10000031B11	422	EFA100190	4884	EFA1c0041_orf_56p	10480
E31V110000031B12	423	EFA100042	4004	EFA160041_0H_30P	10/92

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
E3M10000031C01	424	EFA102736	5007	EFA1c0022_orf_60p	10556
E3M10000031C04	425	EFA101414	4939	EFA1c0022_orf_15p	10528
E3M10000031C06	426	EFA100704	4887	EFA1c0010_orf_4p	10482
E3M10000031C10	427	EFA101411	4936	EFA1c0022_orf_13p	10526
E3M10000031C11	428	EFA101120	4911	EFA1c0036 orf 113p	10687
E3M10000031C12	429	EFA100668	4885	EFA1c0035 orf 58p	10679
E3M10000031D03	430	EFA102503	4996	EFA1c0032 orf 32p	10643
E3M10000031D04	431	EFA102502	4995	EFA1c0031 orf 36p	10627
E3M10000031D08	432	EFA102503	4996	EFA1c0032 orf 32p	10643
E3M10000031E03	433	EFA102501	4994	EFA1c0031 orf 35p	10626
E3M10000031E09	434	EFA102736	5007	EFA1c0022_orf_60p	10556
E3M10000031F02	435	EFA100642	4884	EFA1c0041 orf 56p	10792
E3M10000031F02	435	EFA101685	4952	EFA1c0041 orf 55p	10791
E3M10000031F04	436	EFA101160	4917	EFA1c0022 orf 3p	10549
E3M10000031F07	437	EFA102656	5004	EFA1c0039 orf 26p	10734
E3M10000031F09	438	EFA102764	5008	EFA1c0008 orf 3p	10478
E3M10000031F11	439	EFA102549	5000	EFA1c0022 orf 24p	10538
E3M10000031F11	439	EFA102551	5001	EFA1c0022 orf 25p	10539
E3M10000031G03	440	EFA102655	5003	EFA1c0039 orf 25p	10733
E3M10000031G04	441	EFA103571	5030	EFA1c0044 orf 101p	10879
E3M10000031G05	442	EFA102501	4994	EFA1c0031 orf 35p	10626
E3M10000031G06	443	EFA102656	5004	EFA1c0039 orf 26p	10734
E3M10000031G07	444	EFA103038	5017	EFA1c0030 orf 17p	10613
E3M10000031G08	445	EFA100295	4873	EFA1c0021 orf 15p	10517
E3M10000031G11	446	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000031H05	447	EFA101121	4912	EFA1c0036 orf 112p	10686
E3M10000031H06	448	EFA101540	4947	EFA1c0012 orf 4p	10487
E3M10000031H07	449	EFA103038	5017	EFA1c0030 orf 17p	10613
E3M10000031H08	450	EFA102736	5007	EFA1c0022 orf 60p	10556
E3M10000031H10	451	EFA103038	5017	EFA1c0030 orf 17p	10613
E3M10000031H11	452	EFA100642	4884	EFA1c0041 orf 56p	10792
E3M10000031H11	452	EFA101685	4952	EFA1c0041 orf 55p	10791
E3M10000032A02	453	EFA100704	4887	EFA1c0010 orf 4p	10482
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E3M10000032A06	455	EFA101022	4906	EFA1c0043 orf 69p	10875
E3M10000032A07	456	EFA101670	4950	EFA1c0019_orf_20p	10511
E3M10000032A08	457	EFA100329	4875	EFA1c0041 orf 35p	10782
E3M10000032A09	458	EFA100394	4876	EFA1c0034 orf 6p	10675
E3M10000032A10	459	EFA101410	4935	EFA1c0022 orf 12p	10525
E3M10000032A11	460	EFA100642	4884	EFA1c0041 orf 56p	10792
E3M10000032A11	460	EFA101685	4952	EFA1c0041_orf_55p	10792
E3M10000032R11	461	EFA101540	4947	EFA1c0041_off_33p	10791
E3M10000032B04	462	EFA102091	4947	EFA1c0012 orf 3p	10487
E3M10000032B07	463	EFA101164	4921	EFA1c0022_orf_7p	10558
E3M10000032B08	464	EFA102698	5005	EFA1c0045_orf_115p	10909
E3M10000032B08	465	EFA102051	4976	#N/A	#N/A
E3M10000032B09	466	EFA102091	4977	EFA1c0010 orf 3p	#N/A 10481
E3M10000032B11	467	EFA102091	4873	EFA1c0010_orf_3p	10481
E3M10000032B12	407	EFA100293	48/3	EFAICUUZI_OTI_ISP	10317

Clone name	Clone SeqID	PathoSeq Locus	Gene SegID (protein)	Genemarked gene	full length ORF Protein Seq ID
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E3M10000032C02	469	EFA100151	4864	EFA1c0021_orf_14p	10516
E3M10000032C03	470	EFA103348	5025	EFA1c0043_orf_67p	10873
E3M10000032C04	471	EFA101163	4920	EFA1c0022 orf 6p	10557
E3M10000032C06	472	EFA101150	4915	EFA1c0038 orf 57p	10719
E3M10000032C09	473	EFA100740	4889	EFA1c0022 orf 22p	10536
E3M10000032C11	474	EFA102501	4994	EFA1c0031_orf_35p	10626
E3M10000032C12	475	EFA101165	4922	EFA1c0022 orf 8p	10559
E3M10000032D01	476	EFA103504	5028	EFA1c0033 orf 94p	10671
E3M10000032D02	477	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000032D03	478	EFA100399	4878	EFA1c0041_orf_104p	10766
E3M10000032D06	479	EFA100151	4864	EFA1c0021_orf_14p	10516
E3M10000032D09	480	EFA100151	4864	EFA1c0021 orf 14p	10516
E3M10000032D12	481	EFA101165	4922	EFA1c0022 orf 8p	10559
E3M10000032E04	482	EFA101792	4961	EFA1c0042 orf 113p	10805
E3M10000032E04	482	EFA103786	5031	EFA1c0042 orf 114p	10806
E3M10000032E05	483	EFA103038	5017	EFA1c0030 orf 17p	10613
E3M10000032E08	484	EFA101164	4921	EFA1c0022_orf 7p	10558
E3M10000032E10	485	EFA100870	4899	EFA1c0031 orf 36p	10627
E3M10000032E10	485	EFA102502	4995	EFA1c0031 orf 36p	10627
E3M10000032E11	486	EFA101414	4939	EFA1c0022 orf 15p	10528
E3M10000032E12	487	EFA102326	4986	#N/A	#N/A
E3M10000032F02	488	EFA100210	4870	EFA1c0022 orf 9p	10560
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E3M10000032F03	489	EFA101414	4939	EFA1c0022 orf 15p	10528
E3M10000032F05	490	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000032F07	491	EFA102780	5010	EFA1c0045_orf_101p	10908
E3M10000032F08	492	EFA102501	4994	EFA1c0031 orf 35p	10626
E3M10000032F11	493	EFA100642	4884	EFA1c0041 orf 56p	10792
E3M10000032F12	494	EFA102201	4982	#N/A	#N/A
E3M10000032G01	495	EFA102502	4995	EFA1c0031 orf 36p	10627
E3M10000032G02	496	EFA100870	4899	EFA1c0031_orf_36p	10627
E3M10000032G04	497	EFA100704	4887	EFA1c0010_orf_4p	10482
E3M10000032G05	498	EFA101540	4947	EFA1c0012_orf_4p	10487
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E3M10000033A03	506	EFA101253	4924	EFA1c0043 orf 178p	10852
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E3M10000033A05	508	EFA102551	5001	EFA1c0022 orf 25p	10539
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E3M10000033A08	511	EFA102656	5004	EFA1c0039_orf_26p	10734
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Clone name	Clone SeqID	PathoSeq Locus	Gene SegID (protein)	Genemarked gene	full length ORF Protein Seq
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E3M10000033B02	514	EFA101412	4937	EFA1c0022_orf_14p	10527
E3M10000033B04	515	EFA101765	4958	EFA1c0025_orf_33p	10587
E3M10000033B05	516	EFA102541	4998	EFA1c0028_orf_3p	10602
E3M10000033B06	517	EFA102351	4989	EFA1c0032_orf_20p	10634
E3M10000033B08	518	EFA102091	4977	EFA1c0010 orf 3p	10481
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E3M10000033C02	521	EFA103174	5021	EFA1c0036 orf 120p	10689
E3M10000033C05	522	EFA102541	4998	EFA1c0028 orf 3p	10602
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E3M10000033C10	524	EFA101411	4936	EFA1c0022 orf 13p	10526
E3M10000033C11	525	EFA103504	5028	EFA1c0033 orf 94p	10671
E3M10000033C12	526	EFA102389	4992	EFA1c0044 orf 83p	10904
E3M10000033D01	527	EFA102351	4989	EFA1c0032 orf 20p	10634
E3M10000033D04	528	EFA101682	4951	EFA1c0041 orf 53p	10789
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E3M10000033D06	530	EFA100641	4883	EFA1c0041 orf 57p	10793
E3M10000033D06	530	EFA100642	4884	EFA1c0041_orf_56p	10792
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B3MI0000033G08	Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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E3MI0000033H04 560	E3M10000033G12	558	EFA101686	4953	EFA1c0045_orf_63p	10940
E3M10000033H05 S61	E3M10000033H02	559	EFA101415	4940	EFA1c0022_orf_16p	10529
E3MI0000033H07 562 EFA102502 4995 EFA10031_ort_36p 10627 E3MI0000033H08 563 EFA101160 4917 EFA10022_ort_3p 10549 E3MI0000033H09 565 EFA101079 4988 #FA10001_ort_56p 10792 E3MI0000033H10 565 EFA100190 4867 EFA10010_ort_2p 10480 E3MI0000034A02 567 EFA102501 4994 EFA10032_ort_37p 10542 E3MI0000034A03 568 EFA103338 5017 EFA10033_ort_17p 10541 E3MI0000034A04 569 EFA103504 5028 EFA10033_ort_17p 10648 E3MI0000034B02 570 EFA103504 5028 EFA10033_ort_19p 10671 E3MI0000034B02 570 EFA102552 4995 EFA10033_ort_29p 10671 E3MI0000034D01 573 EFA100190 4867 EFA100031_ort_36p 10627 E3MI0000034D02 574 EFA100162 4919 EFA100010_ort_2p 10480 E3MI0000034F03 578 EFA101162 <td>E3M10000033H04</td> <td>560</td> <td>EFA102780</td> <td>5010</td> <td>EFA1c0045_orf_101p</td> <td>10908</td>	E3M10000033H04	560	EFA102780	5010	EFA1c0045_orf_101p	10908
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E3M10000034F04	E3M10000034E01	575	EFA101162	4919		10555
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E3M10000035A05 586 EFA101540 4947 EFA1c0012_orf_4p 10487 E3M10000035A06 587 EFA103571 5030 EFA1c0044_orf_101p 10879 E3M10000035A08 588 EFA103038 5017 EFA1c0030_orf_17p 10613 E3M10000035A09 589 EFA100210 4870 EFA1c0022_orf_9p 10560 E3M10000035A11 590 EFA100151 4864 EFA1c0021_orf_14p 10516 E3M10000035B01 591 EFA101022 4906 EFA1c0043_orf_69p 10875 E3M10000035B03 592 EFA100704 4887 EFA1c0010_orf_4p 10482 E3M10000035B06 593 EFA101164 4921 EFA1c0022_orf_7p 10558 E3M10000035B07 594 EFA103571 5030 EFA1c0044_orf_101p 10879 E3M10000035B08 595 EFA102780 5010 EFA1c0044_orf_101p 10908 E3M10000035B11 597 EFA103571 5030 EFA1c0001_orf_14p 10516 E3M10000035B12 598 EF	E3M10000035A04	585	EFA103571	5030		
E3M10000035A06 587 EFA103571 5030 EFA1c0044_orf_101p 10879 E3M10000035A08 588 EFA103038 5017 EFA1c0030_orf_17p 10613 E3M10000035A09 589 EFA100210 4870 EFA1c0022_orf_9p 10560 E3M1000035A11 590 EFA100151 4864 EFA1c0021_orf_14p 10516 E3M1000035B01 591 EFA101022 4906 EFA1c0043_orf_69p 10875 E3M10000035B03 592 EFA100704 4887 EFA1c0010_orf_4p 10482 E3M10000035B06 593 EFA101164 4921 EFA1c0022_orf_7p 10558 E3M10000035B07 594 EFA103571 5030 EFA1c0044_orf_101p 10879 E3M10000035B08 595 EFA102780 5010 EFA1c0045_orf_101p 10908 E3M10000035B1 596 EFA100151 4864 EFA1c0021_orf_14p 10516 E3M10000035B1 596 EFA100151 4864 EFA1c0021_orf_14p 10516 E3M10000035B1 597 EFA103571 5030 EFA1c0044_orf_101p 10879 E3M10000035B1 596 EFA100151 4864 EFA1c0021_orf_14p 10516 E3M10000035B1 597 EFA103571 5030 EFA1c0044_orf_101p 10879 E3M10000035B1 598 EFA10338 5017 EFA1c0030_orf_17p 10613 E3M10000035C03 600 EFA101417 4942 EFA1c0022_orf_18p 10531 E3M10000035C04 601 EFA103038 5017 EFA1c0030_orf_17p 10613 E3M10000035C05 602 EFA100870 4899 EFA1c0031_orf_36p 10627	E3M10000035A05	586	EFA101540	4947		10487
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E3M10000035A11 590 EFA100151 4864 EFA1c0021_orf_14p 10516 E3M10000035B01 591 EFA101022 4906 EFA1c0043_orf_69p 10875 E3M10000035B03 592 EFA100704 4887 EFA1c0010_orf_4p 10482 E3M10000035B06 593 EFA101164 4921 EFA1c0022_orf_7p 10558 E3M10000035B07 594 EFA103571 5030 EFA1c0044_orf_101p 10879 E3M10000035B08 595 EFA102780 5010 EFA1c0045_orf_101p 10908 E3M10000035B10 596 EFA100151 4864 EFA1c0021_orf_14p 10516 E3M10000035B11 597 EFA103571 5030 EFA1c0044_orf_101p 10879 E3M10000035B1 598 EFA103038 5017 EFA1c0030_orf_17p 10613 E3M10000035C01 599 EFA100704 4887 EFA1c0010_orf_4p 10482 E3M10000035C03 600 EFA101417 4942 EFA1c0002_orf_18p 10531 E3M10000035C04 601 EF	E3M10000035A08	588	EFA103038	5017		10613
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E3M10000035C04 601 EFA103038 5017 EFA1c0030_orf_17p 10613 E3M10000035C05 602 EFA100870 4899 EFA1c0031_orf_36p 10627						
E3M10000035C05 602 EFA100870 4899 EFA1c0031_orf_36p 10627					- - .	
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Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq
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E3M10000035C08	605	EFA100742	4891	EFA1c0022 orf 20p	10534
E3M10000035C09	606	EFA103062	5019	EFA1c0030 orf 19p	10615
E3M10000035C11	607	EFA100704	4887	EFA1c0010 orf 4p	10482
E3M10000035C12	608	EFA100704	4887	EFA1c0010 orf 4p	10482
E3M10000035D02	609	EFA101160	4917	EFA1c0022 orf 3p	10549
E3M10000035D03	610	EFA103504	5028	EFA1c0033 orf 94p	10671
E3M10000035D04	611	EFA101540	4947	EFA1c0012 orf 4p	10487
E3M10000035D05	612	EFA101162	4919	EFA1c0022_orf_5p	10555
E3M10000035D10	613	EFA103571	5030	EFA1c0044_orf_101p	10879
E3M10000035D11	614	EFA100919	4901	EFA1c0013 orf 12p	10491
E3M10000035E03	615	EFA101414	4939	EFA1c0022 orf 15p	10528
E3M10000035E04	616	EFA101141	4914	EFA1c0030 orf 18p	10614
E3M10000035E05	617	EFA102006	4973	EFA1c0025 orf 17p	10580
E3M10000035E07	618	EFA100919	4901	EFA1c0013 orf 12p	10491
E3M10000035E08	619	EFA101162	4919	EFA1c0022_orf_5p	10555
E3M10000035E09	620	EFA100312	4874	EFA1c0032_orf_28p	10641
E3M10000035E10	621	EFA101022	4906	EFA1c0043 orf 69p	10875
E3M10000035E11	622	EFA100870	4899	EFA1c0031 orf 36p	10627
E3M10000035E12	623	EFA100704	4887	EFA1c0010 orf 4p	10482
E3M10000035F01	624	EFA101417	4942	EFA1c0022 orf 18p	10531
E3M10000035F02	625	EFA101925	4971	EFA1c0044 orf 19p	10893
E3M10000035F03	626	EFA100312	4874	EFA1c0032 orf 28p	10641
E3M10000035F06	627	EFA101080	4909	#N/A	#N/A
E3M10000035F07	628	EFA101165	4922	EFA1c0022 orf 8p	10559
E3M10000035F08	629	EFA100704	4887	EFA1c0010 orf 4p	10482
E3M10000035F09	630	EFA101410	4935	EFA1c0022 orf 12p	10525
E3M10000035F09	630	EFA101411	4936	EFA1c0022_orf_13p	10526
E3M10000035F11	631	EFA100704	4887	EFA1c0010 orf 4p	10482
E3M10000035F12	632	EFA101120	4911	EFA1c0036 orf 113p	10687
E3M10000035F12	633	EFA100190	4867	EFA1c0010 orf 2p	10480
E3M10000035G02	633	EFA102091	4977	EFA1c0010_orf 3p	10481
E3M10000035G02	634	EFA100210	4870	EFA1c0022_orf_9p	10560
E3M10000035G04	635	EFA102502	4995	EFA1c0031 orf 36p	10627
E3M10000035G08	636	EFA100642	4884	EFA1c0041 orf 56p	10792
E3M10000035G09	637	EFA103504	5028	EFA1c0041_off_50p	10671
E3M10000035G09	637	EFA103504	5029	EFA1c0033_orf_95p	10672
E3M10000035G02	638	EFA103038	5017	EFA1c0030_orf_17p	10613
E3M10000035G10	639	EFA101540	4947	EFA1c0012 orf 4p	10013
E3M10000035H03	640	EFA101080	4909	#N/A	#N/A
E3M10000035H03	641	EFA100210	4870	EFA1c0022 orf 9p	10560
E3M10000035H09	642	EFA102501	4994	EFA1c0022_0f1_9p	10560
E3M10000035H09	643	EFA101257	4925	EFA1c0031_ort_35p EFA1c0045 orf 159p	10026
E3M10000035H11	643	EFA101257 EFA101258	4925	EFA1c0045_orf_159p EFA1c0045_orf_160p	10917
E3M10000035H11	644	EFA101258 EFA103504	5028	EFA1c0045_orf_160p	10918
E3M10000036A03	645	EFA103504 EFA101416	4941		
E31V11UUUUU36AU4	645	EFA101416 EFA102780	5010	EFA1c0022_orf_17p EFA1c0045_orf_101p	10530

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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E3M10000036A07	648	EFA103268	5023	EFA1c0010 orf 1p	10479
E3M10000036A08	649	EFA103038	5017	EFA1c0030 orf 17p	10613
E3M10000036A09	650	EFA101165	4922	EFA1c0022 orf 8p	10559
E3M10000036A10	651	EFA100210	4870	EFA1c0022 orf 9p	10560
E3M10000036B01	652	EFA101121	4912	EFA1c0036 orf 112p	10686
E3M10000036B03	653	EFA101686	4953	EFA1c0045 orf 63p	10940
E3M10000036B06	654	EFA101162	4919	EFA1c0022_orf_5p	10555
E3M10000036B07	655	EFA103038	5017	EFA1c0030 orf 17p	10613
E3M10000036B08	656	EFA100151	4864	EFA1c0021 orf 14p	10516
E3M10000036B09	657	EFA100190	4867	EFA1c0010_orf_2p	10480
E3M10000036B11	658	EFA103504	5028	EFA1c0033_orf_94p	10671
E3M10000036B12	659	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000036B12	659	EFA101163	4920	EFA1c0022_orf_6p	10557
E3M10000036C01	660	EFA101416	4941	EFA1c0022_orf_17p	10537
E3M10000036C03	661	EFA103571	5030	EFA1c0022_011_17p	10330
E3M10000036C05	662	EFA102091	4977	EFA1c0010 orf 3p	10481
E3M10000036C08	663	EFA101141	4914	EFA1c0010_off_3p	10481
E3M1000036C07	664	J]	_ ~ .	
		EFA100151	4864	EFA1c0021_orf_14p	10516
E3M10000036C09	665	EFA101540	4947	EFA1c0012_orf_4p	10487
E3M10000036C10	666	EFA101540	4947	EFA1c0012_orf_4p	10487
E3M10000036C11	667	EFA101417	4942	EFA1c0022_orf_18p	10531
E3M10000036D03	668	EFA100704	4887	EFA1c0010_orf_4p	10482
E3M10000036D04	669	EFA102201	4982	#N/A	#N/A
E3M10000036D06	670	EFA100740	4889	EFA1c0022_orf_22p	10536
E3M10000036D08	671	EFA101164	4921	EFA1c0022_orf_7p	10558
E3M10000036D09	672	EFA103571	5030	EFA1c0044_orf_101p	10879
E3M10000036D10	673	EFA101121	4912	EFA1c0036_orf_112p	10686
E3M10000036D11	674	EFA102502	4995	EFA1c0031_orf_36p	10627
E3M10000036D12	675	EFA102091	4977	EFA1c0010_orf_3p	10481
E3M10000036E01	676	EFA101121	4912	EFA1c0036_orf_112p	10686
E3M10000036E04	677	EFA101162	4919	EFA1c0022_orf_5p	10555
E3M10000036E05	678	EFA101414	4939	EFA1c0022_orf_15p	10528
E3M10000036E07	679	EFA101022	4906	EFA1c0043_orf_69p	10875
E3M10000036E08	680	EFA100210	4870	EFA1c0022_orf_9p	10560
E3M10000036F03	681	EFA100210	4870	EFA1c0022_orf_9p	10560
E3M10000036F04	682	EFA101686	4953	EFA1c0045_orf_63p	10940
E3M10000036F05	683	EFA101792	4961	EFA1c0042_orf_113p	10805
E3M10000036F08	684	EFA102091	4977	EFA1c0010_orf_3p	10481
E3M10000036F09	685	EFA101404	4933	EFA1c0033_orf_55p	10663
E3M10000036F10	686	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000036F12	687	EFA101163	4920	EFA1c0022 orf 6p	10557
E3M10000036G01	688	EFA102549	5000	EFA1c0022 orf 24p	10538
E3M10000036G01	688	EFA102551	5001	EFA1c0022 orf 25p	10539
E3M10000036G02	689	EFA101121	4912	EFA1c0036 orf 112p	10686
E3M10000036G03	690	EFA102656	5004	EFA1c0039 orf 26p	10734
E3M10000036G04	691	EFA102091	4977	EFA1c0010 orf 3p	10481
E3M10000036G06	692	EFA100295	4873	EFA1c0021 orf 15p	10517

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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E3M10000036H02	694	EFA103038	5017	EFA1c0030_orf_17p	10613
E3M10000036H03	695	EFA103571	5030	EFA1c0044_orf_101p	10879
E3M10000036H04	696	EFA103365	5026	EFA1c0022_orf_1p	10533
E3M10000036H05	697	EFA100194	4868	EFA1c0022_orf_26p	10540
E3M10000036H06	698	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000036H07	699	EFA102501	4994	EFA1c0031_orf_35p	10626
E3M10000036H08	700	EFA103210	5022	EFA1c0036 orf 119p	10688
E3M10000036H09	701	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000036H10	702	EFA101141	4914	EFA1c0030_orf_18p	10614
E3M10000037A03	703	EFA102091	4977	EFA1c0010 orf 3p	10481
E3M10000037A06	704	EFA100870	4899	EFA1c0031_orf_36p	10627
E3M10000037A08	705	EFA103365	5026	EFA1c0022 orf 1p	10533
E3M10000037A09	706	EFA100756	4893	EFA1c0024 orf 39p	10575
E3M10000037A10	707	EFA103268	5023	EFA1c0010 orf 1p	10479
E3M10000037B02	708	EFA100641	4883	EFA1c0041 orf 57p	10793
E3M10000037B02	708	EFA100642	4884	EFA1c0041 orf 56p	10792
E3M10000037B07	709	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000037B08	710	EFA100151	4864	EFA1c0021 orf 14p	10516
E3M10000037B11	711	EFA101686	4953	EFA1c0045_orf_63p	10940
E3M10000037C01	712	EFA101080	4909	#N/A	#N/A
E3M10000037C02	713	EFA102351	4989	EFA1c0032 orf 20p	10634
E3M10000037C04	714	EFA103504	5028	EFA1c0033 orf 94p	10671
E3M10000037C05	715	EFA102655	5003	EFA1c0039 orf 25p	10733
E3M10000037C07	716	EFA101160	4917	EFA1c0022 orf 3p	10549
E3M10000037C07	716	EFA101161	4918	EFA1c0022 orf 4p	10551
E3M10000037C11	717	EFA100615	4881	EFA1c0016 orf 29p	10501
E3M10000037C12	718	EFA102502	4995	EFA1c0031 orf 36p	10627
E3M10000037D02	719	EFA100642	4884	EFA1c0041 orf 56p	10792
E3M10000037D03	720	EFA100795	4896	EFA1c0043 orf 229p	10863
E3M10000037D03	720	EFA103081	5020	EFA1c0043 orf 228p	10862
E3M10000037D04	721	EFA100210	4870	EFA1c0022 orf 9p	10560
E3M10000037D05	722	EFA101416	4941	EFA1c0022 orf 17p	10530
E3M10000037D06	723	EFA101161	4918	EFA1c0022 orf 4p	10551
E3M10000037D09	724	EFA100190	4867	EFA1c0010 orf 2p	10480
E3M10000037D09	724	EFA102091	4977	EFA1c0010 orf 3p	10481
E3M10000037D11	725	EFA100210	4870	EFA1c0022 orf 9p	10560
E3M10000037E01	726	EFA102736	5007	EFA1c0022 orf 60p	10556
E3M10000037E02	727	EFA100704	4887	EFA1c0010_orf_4p	10482
E3M10000037E03	728	EFA102503	4996	EFA1c0032 orf 32p	10643
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E3M10000037E07	730	EFA100210	4870	EFA1c0022 orf 9p	10560
E3M10000037E08	731	EFA100642	4884	EFA1c0041 orf 56p	10792
E3M10000037E10	732	EFA101253	4924	EFA1c0043 orf 178p	10852
E3M10000037E12	733	EFA101686	4953	EFA1c0045_orf_63p	10940
E3M10000037E12	734	EFA103504	5028	EFA1c0043_orf_94p	10940
E3M10000037F02	735	EFA101160	4917	EFA1c0022 orf 3p	10549
E3M10000037F06	736	EFA100210	4870	EFA1c0022_orf_9p	10560

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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E3M10000037G02	740	EFA101165	4922	EFA1c0022 orf 8p	10559
E3M10000037G03	741	EFA102780	5010	EFA1c0045 orf 101p	10908
E3M10000037G05	742	EFA102780	5010	EFA1c0045 orf 101p	10908
E3M10000037G06	743	EFA103295	5024	EFA1c0032 orf 1p	10633
E3M10000037G07	744	EFA101541	4948	EFA1c0012 orf 5p	10488
E3M10000037G08	745	EFA101121	4912	EFA1c0036 orf 112p	10686
E3M10000037G10	746	EFA101412	4937	EFA1c0022 orf 14p	10527
E3M10000037G11	747	EFA103038	5017	EFA1c0030 orf 17p	10613
E3M10000037H02	748	EFA101413	4938	#N/A	#N/A
E3M10000037H05	749	EFA101686	4953	EFA1c0045 orf 63p	10940
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E3M10000037H11	752	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000038A02	753	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000038A03	754	EFA103038	5017	EFA1c0030 orf 17p	10613
E3M10000038A05	755	EFA100151	4864	EFA1c0030_011_17p	10516
E3M10000038A06	756	EFA102549	5000	EFA1c0022 orf 24p	10538
E3M10000038A07	757	EFA102501	4994	EFA1c0031 orf 35p	10626
E3M10000038A09	758	EFA102736	5007	EFA1c0031_011_53p	10556
E3M10000038A10	759	EFA100210	4870	EFA1c0022_orf_9p	10550
E3M10000038A11	760	EFA101417	4942	EFA1c0022_0ff_9p	10531
E3M10000038A11	761	EFA103210	5022	EFA1c0022_0ff_18p	10688
E3M10000038B02	762	EFA102389	4992	EFA1c0034_orf_83p	10904
E3M10000038B03	763	EFA101414	4939	EFA1c0022 orf 15p	10528
E3M10000038B05	764	EFA100795	4896	EFA1c0022_0ff_15p	10328
E3M10000038B05	764	EFA103081	5020	EFA1c0043_orf_228p	10862
E3M10000038B07	765	EFA100190	4867	EFA1c0010 orf 2p	10480
E3M10000038B07	766	EFA101160	4917	EFA1c0022 orf 3p	10480
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E3M10000038B09	768	EFA102656	5004	EFA1c0039 orf 26p	10734
E3M10000038C02	769	EFA102780	5010	EFA1c0039_0ff_20p	10734
E3M10000038C02	770	EFA102656	5004	EFA1c0039 orf 26p	10734
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E3M10000038C03	772	EFA101963	4972	EFA1c0043_orf_162p	10940
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				EFA1c0012_orf_4p	
E3M10000038D08	778	EFA101160	4917	EFA1c0022_orf_3p	10549
E3M10000038D10	779	EFA103504	5028	EFA1c0033_orf_94p	10671
E3M10000038D11	780	EFA103571	5030	EFA1c0044_orf_101p	10879
E3M10000038D12	781	EFA101540	4947	EFA1c0012_orf_4p	10487
E3M10000038E02	782	EFA100704	4887	EFA1c0010_orf_4p	10482
E3M10000038E03	783	EFA101159	4916	EFA1c0022_orf_2p	10543

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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E3M10000038E08	787	EFA101121	4912	EFA1c0036 orf 112p	10686
E3M10000038E11	788	EFA102780	5010	EFA1c0045_orf_101p	10908
E3M10000038F02	789	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000038F04	790	EFA101686	4953	EFA1c0045 orf 63p	10940
E3M10000038F05	791	EFA101160	4917	EFA1c0022 orf 3p	10549
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E3M10000038F06	792	EFA103571	5030	EFA1c0044 orf 101p	10879
E3M10000038F07	793	EFA103210	5022	EFA1c0036_orf_119p	10688
E3M10000038F09	794	EFA102185	4980	EFA1c0045 orf 95p	10950
E3M10000038F10	795	EFA101080	4909	#N/A	#N/A
E3M10000038F11	796	EFA100740	4889	EFA1c0022 orf 22p	10536
E3M10000038G02	797	EFA100919	4901	EFA1c0013 orf 12p	10491
E3M10000038G03	798	EFA101414	4939	EFA1c0022 orf 15p	10528
E3M10000038G06	799	EFA100704	4887	EFA1c0010 orf 4p	10482
E3M10000038G07	800	EFA102352	4990	EFA1c0032 orf 21p	10635
E3M10000038G07	800	EFA102353	4991	EFA1c0032 orf 22p	10636
E3M10000038G11	801	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000038H02	802	EFA101414	4939	EFA1c0022 orf 15p	10528
E3M10000038H05	803	EFA101160	4917	EFA1c0022 orf 3p	10549
E3M10000038H06	804	EFA100295	4873	EFA1c0021 orf 15p	10517
E3M10000038H07	805	EFA101417	4942	EFA1c0022 orf 18p	10531
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E3M10000038H09	807	EFA102802	5012	EFA1c0043 orf 18p	10854
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E3M10000039A02	809	EFA101737	4956	EFA1c0041 orf 15p	10778
E3M10000039A06	810	EFA102656	5004	EFA1c0039 orf 26p	10734
E3M10000039A07	811	EFA102006	4973	EFA1c0025 orf 17p	10580
E3M10000039A08	812	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000039A10	813	EFA101257	4925	EFA1c0045 orf 159p	10917
E3M10000039A11	814	EFA101412	4937	EFA1c0022 orf 14p	10527
E3M10000039B01	815	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000039B03	816	EFA101160	4917	EFA1c0022 orf 3p	10549
E3M10000039B04	817	EFA101415	4940	EFA1c0022 orf 16p	10529
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E3M10000039B06	818	EFA100870	4899	EFA1c0031 orf 36p	10627
E3M10000039B07	819	EFA102110	4978	EFA1c0042_orf_99p	10841
E3M10000039B08	820	EFA101416	4941	EFA1c0022_orf 17p	10530
E3M10000039B09	821	EFA101792	4961	EFA1c0042_orf_113p	10805
E3M10000039B11	822	EFA101080	4909	#N/A	#N/A
E3M10000039C02	823	EFA103062	5019	EFA1c0030 orf 19p	10615
E3M10000039C04	824	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000039C04	825	EFA100739	4888	EFA1c0022_orf_23p	10537
E3M10000039C05	826	EFA103504	5028	EFA1c0033 orf 94p	10537
E3M10000039C00	827	EFA101791	4960	EFA1c0042 orf 112p	10804
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Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
E3M10000039C07	827	EFA101792	4961	EFA1c0042_orf_113p	10805
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E3M10000039C09	829	EFA102503	4996	EFA1c0032_orf_32p	10643
E3M10000039C10	830	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000039D02	831	EFA101165	4922	EFA1c0022 orf 8p	10559
E3M10000039D03	832	EFA102655	5003	EFA1c0039 orf 25p	10733
E3M10000039D04	833	EFA102656	5004	EFA1c0039 orf 26p	10734
E3M10000039D06	834	EFA101540	4947	EFA1c0012_orf_4p	10487
E3M10000039E01	835	EFA102201	4982	#N/A	#N/A
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E3M10000039E03	837	EFA100919	4901	EFA1c0013_orf_12p	10491
E3M10000039E05	838	EFA101686	4953	EFA1c0045_orf_63p	10940
E3M10000039E07	839	EFA103295	5024	EFA1c0032 orf 1p	10633
E3M10000039E08	840	EFA101685	4952	EFA1c0041 orf 55p	10791
E3M10000039F01	841	EFA102656	5004	EFA1c0039 orf 26p	10734
E3M10000039F02	842	EFA103021	5015	EFA1c0030_orf_16p	10612
E3M10000039F03	843	EFA102788	5011	EFA1c0033 orf 41p	10661
E3M10000039F03	843	EFA103375	5027	EFA1c0033 orf 40p	10660
E3M10000039F06	844	EFA100739	4888	EFA1c0022 orf 23p	10537
E3M10000039F07	845	EFA102541	4998	EFA1c0028_orf_3p	10602
E3M10000039F08	846	EFA101162	4919	EFA1c0022 orf 5p	10555
E3M10000039G01	847	EFA102502	4995	EFA1c0031 orf 36p	10627
E3M10000039G02	848	EFA101686	4953	EFA1c0045 orf 63p	10940
E3M10000039G05	849	EFA100919	4901	EFA1c0013 orf 12p	10491
E3M10000039G07	850	EFA101686	4953	EFA1c0045 orf 63p	10940
E3M10000039G09	851	EFA102541	4998	EFA1c0028 orf 3p	10602
E3M10000039G10	852	EFA101682	4951	EFA1c0041 orf 53p	10789
E3M10000039H02	853	EFA101160	4917	EFA1c0022 orf 3p	10549
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E3M10000040A03	858	EFA101123	4913	EFA1c0040 orf 22p	10748
E3M10000040A05	859	EFA101080	4909	#N/A	#N/A
E3M10000040A07	860	EFA100157	4865	EFA1c0034 orf 63p	10673
E3M10000040A09	861	EFA102502	4995	EFA1c0031 orf 36p	10627
E3M10000040A10	862	EFA101417	4942	EFA1c0022 orf 18p	10531
E3M10000040A11	863	EFA101685	4952	EFA1c0041 orf 55p	10791
E3M10000040B01	864	EFA102788	5011	EFA1c0033 orf 41p	10661
E3M10000040B02	865	EFA102655	5003	EFA1c0039_orf_25p	10733
E3M10000040B05	866	EFA100190	4867	EFA1c0010 orf 2p	10480
E3M10000040B05	866	EFA103268	5023	EFA1c0010_orf_lp	10480
E3M10000040B06	867	EFA102518	4997	EFA1c0032 orf 46p	10479
E3M10000040B08	868	EFA100919	4901	EFA1c0032_011_40p	10491
E3M10000040B08	869	EFA102502	4995	EFA1c0013_orf_12p	10491
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E3M10000040B10	870	EFA102656	5004	EFA1c0039_orf_26p	10734

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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E3M10000040C05	874	EFA102501	4994	EFA1c0031_orf_35p	10626
E3M10000040C06	875	EFA102091	4977	EFA1c0010_orf_3p	10481
E3M10000040C07	876	EFA101121	4912	EFA1c0036 orf 112p	10686
E3M10000040C08	877	EFA102780	5010	EFA1c0045_orf_101p	10908
E3M10000040C09	878	EFA100165	4866	EFA1c0032 orf 23p	10637
E3M10000040C09	878	EFA102353	4991	EFA1c0032 orf 22p	10636
E3M10000040C10	879	EFA101686	4953	EFA1c0045_orf_63p	10940
E3M10000040C11	880	EFA102501	4994	EFA1c0031 orf 35p	10626
E3M10000040C12	881	EFA102780	5010	EFA1c0045 orf 101p	10908
E3M10000040D03	882	EFA102201	4982	#N/A	#N/A
E3M10000040D04	883	EFA101080	4909	#N/A	#N/A
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E3M10000040D12	885	EFA101686	4953	EFA1c0045 orf 63p	10940
E3M10000040E02	886	EFA102051	4976	#N/A	#N/A
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E3M10000040E11	888	EFA103039	5018	EFA1c0043_orf_16p	10850
E3M10000040E12	889	EFA102091	4977	EFA1c0010 orf 3p	10481
E3M10000040F01	890	EFA100295	4873	EFA1c0021_orf_15p	10517
E3M10000040F03	891	EFA102503	4996	EFA1c0032 orf 32p	10643
E3M10000040F08	892	EFA101080	4909	#N/A	#N/A
E3M10000040F09	893	EFA100919	4901	EFA1c0013 orf 12p	10491
E3M10000040F10	894	EFA102051	4976	#N/A	#N/A
E3M10000040F10	895	EFA101415	4940	EFA1c0022 orf 16p	10529
E3M10000040G02	896	EFA101424	4943	EFA1c0041 orf 39p	10784
E3M10000040G02	896	EFA101425	4944	EFA1c0041_orf_40p	10785
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E3M10000040G05	898	EFA101159	4916	EFA1c0030_0ff_18p	10543
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E3M10000040H05	906	EFA100642	4884	EFA1c0041_orf_56p	10792
E3M10000040H05	906	EFA101685	4952	EFA1c0041_orf_55p	10791
E3M10000040H09	907	EFA101416	4941	EFA1c0022_orf_17p	10530
E3M10000040H09	907	EFA101417	4942	EFA1c0022_orf_18p	10531
E3M10000041A03	908	EFA100615	4881	EFA1c0016_orf_29p	10501
E3M10000041A05	909	EFA102502	4995	EFA1c0031_orf_36p	10627
E3M10000041A08	910	EFA100704	4887	EFA1c0010_orf_4p	10482
E3M10000041A09	911	EFA101354	4930	EFA1c0032_orf_69p	10648
E3M10000041A10	912	EFA100001	4861	EFA1c0030_orf_3p	10618
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Clone name	Clone SegID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF
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E3M10000041B03	915	EFA101414	4939	EFA1c0022_orf_15p	10528
E3M10000041B05	916	EFA102091	4977	EFA1c0010_orf_3p	10481
E3M10000041B06	917	EFA102091	4977	EFA1c0010_orf_3p	10481
E3M10000041B08	918	EFA102655	5003	EFA1c0039_orf_25p	10733
E3M10000041B09	919	EFA101924	4970	EFA1c0044_orf_18p	10891
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E3M10000041B10	920	EFA101080	4909	#N/A	#N/A
E3M10000041B11	921	EFA101416	4941	EFA1c0022_orf_17p	10530
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E3M10000041B12	922	EFA101411	4936	EFA1c0022_orf_13p	10526
E3M10000041C01	923	EFA100151	4864	EFA1c0021_orf_14p	10516
E3M10000041C07	924	EFA100739	4888	EFA1c0022_orf_23p	10537
E3M10000041C08	925	EFA101121	4912	EFA1c0036_orf_112p	10686
E3M10000041C09	926	EFA103365	5026	EFA1c0022_orf_lp	10533
E3M10000041C10	927	EFA102503	4996	EFA1c0032_orf_32p	10643
E3M10000041C11	928	EFA102655	5003	EFA1c0039_orf_25p	10733
E3M10000041C12	929	EFA100798	4897	EFA1c0042_orf_160p	10818
E3M10000041D02	930	EFA102502	4995	EFA1c0031_orf_36p	10627
E3M10000041D03	931	EFA101060	4907	EFA1c0038_orf_73p	10722
E3M10000041D04	932	EFA100642	4884	EFA1c0041_orf_56p	10792
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E3M10000041D11	938	EFA100704	4887	EFA1c0010_orf_4p	10482
E3M10000041D12	939	EFA100394	4876	EFA1c0034_orf_6p	10675
E3M10000041E02	940	EFA101797	4963	EFA1c0045_orf_167p	10924
E3M10000041E03	941	EFA102091	4977	EFA1c0010_orf_3p	10481
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E3M10000041E07	943	EFA102091	4977	EFA1c0010_orf_3p	10481
E3M10000041E10	944	EFA100704	4887	EFA1c0010_orf_4p	10482
E3M10000041E11	945	EFA100190	4867	EFA1c0010_orf_2p	10480
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E3M10000041F06	948	EFA102501	4994	EFA1c0031_orf_35p	10626
E3M10000041F07	949	EFA101159	4916	EFA1c0022_orf_2p	10543
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E3M10000041G02	954	EFA101141	4914	EFA1c0030_orf_18p	10614
E3M10000041G03	955	EFA102253	4984	EFA1c0038_orf_85p	10727
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E3M10000041G08	959	EFA100704	4887	EFA1c0010_orf_4p	10482
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E3M10000041G10	961	EFA100394	4876	EFA1c0034_orf_6p	10675
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E3M10000041H06	965	EFA102502	4995	EFA1c0031 orf 36p	10627
E3M10000041H07	966	EFA103062	5019	EFA1c0030 orf 19p	10615
E3M10000041H08	967	EFA101686	4953	EFA1c0045 orf 63p	10940
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E3M10000041H10	969	EFA101685	4952	EFA1c0041 orf 55p	10791
E3M10000041H111	970	EFA102253	4984	EFA1c0038 orf 85p	10727
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E3M10000042B01	974	EFA101404	4933	EFA1c0033 orf 55p	10663
E3M10000042B02	975	EFA100668	4885	EFA1c0035 orf 58p	10679
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E3M10000042G08	998	EFA102780	5010	EFA1c0045_orf_101p	10908
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E3M10000042G11	999	EFA101121	4912	EFA1c0036_orf_112p	10686
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E3M10000043A02	1004	EFA101799	4964	EFA1c0045_orf_169p	10926
E3M10000043A03	1005	EFA101414	4939	EFA1c0022_orf_15p	10528
E3M10000043A05	1006	EFA102502	4995	EFA1c0031 orf 36p	10627
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E3M10000043B03	1013	EFA103038	5017	EFA1c0030 orf 17p	10613
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E3M10000043B08	1015	EFA101123	4913	EFA1c0040 orf 22p	10748
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E3M10000043B11	1018	EFA100704	4887	EFA1c0010 orf 4p	10482
E3M10000043B12	1019	EFA100151	4864	EFA1c0021 orf 14p	10516
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E3M10000043H08	1049	EFA100615	4881	EFA1c0016_orf_29p	10501
E3M10000043H09	1050	EFA102006	4973	EFA1c0025_orf_17p	10580
E3M10000043H11	1051	EFA102655	5003	EFA1c0039_orf_25p	10733
E3M10000044C02	1052	EFA100955	4902	EFA1c0022_orf_28p	10542
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K1M10000003C01	1055	KPN103882	5040	KPN1c2848_orf_lp	11716
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K1M10000020B02	1065	KPN101729	5036	KPN1c1566 orf 1p	11647
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P1M10000053C02	1177	PA0353	5061	#N/A	#N/A
P1M10000053E07	1178	PA4254	5170	#N/A	#N/A
P1M10000053F08	1179	PA1270	5082	#N/A	#N/A
P1M10000055A11	1180	PA5076	5204	#N/A	#N/A
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P1M10000056C07	1183	PA1360	5084	#N/A	#N/A
P1M10000056F05	1184	PA4258	5173	#N/A	#N/A
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P1M10000056G01	1186	PA5076	5204	#N/A	#N/A
P1M10000058B07	1187	PA5436	5215	#N/A	#N/A
P1M10000059B04	1188	PA4375	5186	#N/A	#N/A
P1M10000059B10	1189	PA4269	5179	#N/A	#N/A
P1M10000059B10	1190	PA0934	5077	#N/A	#N/A
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P1M10000059H08	1191	PA4027	5149	#N/A	#N/A
P1M10000059H09	1192	PA4271	5180	#N/A #N/A	#N/A
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P1M10000061B04	1197	PA2726	5119	#N/A	#N/A
P1M10000061E04	1198	PA4244	5160	#N/A	#N/A
P1M10000061F04	1199	PA3522	5136	#N/A	#N/A
P1M10000062A12	1200	PA4598	5194	#N/A	#N/A
P1M10000062C03	1201	PA0321	5059	#N/A	#N/A
P1M10000062C04	1202	PA4254	5170	#N/A	#N/A
P1M10000062C07	1203	PA4251	5167	#N/A	#N/A
P1M10000062C12	1204	PA5316	5212	#N/A	#N/A
P1M10000062D07	1205	PA4247	5163	#N/A	#N/A
P1M10000062D08	1206	PA0882	5076	#N/A	#N/A
P1M10000062E08	1207	PA4248	5164	#N/A	#N/A
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P1M10000062F06	1208	PA0028	5053	#N/A	#N/A
P1M10000062G11	1209	PA4506	5190	#N/A	#N/A
P1M10000062H01	1210	PA3121	5127	#N/A	#N/A
P1M10000062H04	1211	PA4254	5170	#N/A	#N/A
P1M10000063F02	1212	PA2684	5118	#N/A	#N/A
P1M10000063G02	1213	PA4262	5175	#N/A	#N/A
P1M10000063H02	1214	PA4081	5153	#N/A	#N/A
P1M10000064A10	1215	PA4268	5178	#N/A	#N/A
P1M10000064C02	1216	PA0650	5073	#N/A	#N/A
P1M10000064C03	1217	PA5030	5203	#N/A	#N/A
P1M10000064D03	1218	PA0129	5055	#N/A	#N/A
P1M10000064E05	1219	PA4512	5191	#N/A	#N/A
P1M10000064G12	1220	PA2147	5101	#N/A	#N/A
P1M10000064H07	1221	PA1072	5080	#N/A	#N/A
PIMI000065A04	1222	PA3522	5136	#N/A	#N/A
P1M10000065B07	1223	PA4347	5184	#N/A	#N/A
P1M10000065C03	1224	PA4347	5184	#N/A	#N/A
P1M10000065C05	1225	PA0642	5072	#N/A	#N/A
P1M10000065D06	1226	PA4347	5184	#N/A	#N/A
P1M10000065F01	1227	PA2494	5111	#N/A	#N/A
P1M10000065G06	1228	PA0423	5067	#N/A	#N/A
P1M10000065H07	1229	PA1019	5079	#N/A	#N/A
P1M10000066A10	1230	PA4709	5197	#N/A	#N/A
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P1M10000066F04	1232	PA4024	5148	#N/A	#N/A
P1M10000067A05	1233	PA3876	5144	#N/A	#N/A
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PIM10000067A08	1235	PA0600	5071	#N/A	#N/A
P1M10000067C04	1236	PA3845	5142	#N/A	#N/A
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P1M10000067D05	1238	PA3479	5134	#N/A	#N/A
P1M10000067F05	1239	PA3643	5137	#N/A	#N/A
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PIM1000068A09	1241	PA0353	5061	#N/A	#N/A

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P1M10000068F04	1243	PA4237	5158	#N/A	#N/A
P1M10000068F08	1244	PA5193	5206	#N/A	#N/A
P1M10000068G01	1245	PA3716	5140	#N/A	#N/A
P1M10000068H05	1246	PA4268	5178	#N/A	#N/A
P1M10000069D09	1247	PA4246	5162	#N/A	#N/A
P1M10000069G06	1248	PA4246	5162	#N/A	#N/A
P1M10000069H02	1249	PA4433	5188	#N/A	#N/A
P1M10000070A05	1250	PA2470	5109	#N/A	#N/A
P1M10000070B10	1251	PA5393	5214	#N/A	#N/A
P1M10000070C06	1252	PA4237	5158	#N/A	#N/A
P1M10000070D08	1253	PA4105	5154	#N/A	#N/A
P1M10000070E03	1254	PA4709	5197	#N/A	#N/A
P1M10000070G06	1255	PA3374	5133	#N/A	#N/A
P1M10000070G12	1256	PA3121	5127	#N/A	#N/A
P1M10000070H06	1257	PA3374	5133	#N/A	#N/A
P1M10000071A03	1258	PA4251	5167	#N/A	#N/A
P1M10000071C01	1259	PA4251	5167	#N/A	#N/A
P1M10000071E04	1260	PA3484	5135	#N/A	#N/A
P1M10000071F01	1261	PA0506	5070	#N/A	#N/A
P1M10000073A06	1262	PA4246	5162	#N/A	#N/A
P1M10000073B10	1263	PA5248	5210	#N/A	#N/A
P1M10000073D04	1264	PA1115	5081	#N/A	#N/A
P1M10000073D09	1265	PA1918	5094	#N/A	#N/A
P1M10000073G03	1266	PA5248	5210	#N/A	#N/A
P1M10000074B01	1267	PA4771	5199	#N/A	#N/A
P1M10000074B04	1268	PA1684	5091	#N/A	#N/A
P1M10000074E04	1269	PA0120	5054	#N/A	#N/A
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P1M10000075A04	1273	PA3279	5131	#N/A	#N/A
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P1M10000075B03	1274	PA4576	5193	#N/A	#N/A
P1M10000075F02	1275	PA4254	5170	#N/A	#N/A
P1M10000075G05	1276	PA3709	5139	#N/A	#N/A
P1M10000076D05	1277	PA1876	5093	#N/A	#N/A
P1M10000076D10	1278	PA1636	5090	#N/A	#N/A
P1M10000077A08	1279	PA3479	5134	#N/A	#N/A
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P1M10000079B10	1284	PA4576	5193	#N/A	#N/A
P1M10000079C10	1285	PA4576	5193	#N/A	#N/A
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Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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P1M10000080B01	1289	PA3866	5143	#N/A	#N/A
P1M10000080B06	1290	PA4244	5160	#N/A	#N/A
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P1M10000080C01	1291	PA0469	5068	#N/A	#N/A
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P1M10000080E04	1293	PA4250	5166	#N/A	#N/A
P1M10000081D12	1294	PA3006	5121	#N/A	#N/A
P1M10000081G05	1295	PA4037	5150	#N/A	#N/A
P1M10000081H05	1296	PA4316	5182	#N/A	#N/A
P1M10000082A05	1297	PA0401	5063	#N/A	#N/A
P1M10000082B04	1298	PA3006	5121	#N/A	#N/A
P1M10000082C05	1299	PA4246	5162	#N/A	#N/A
P1M10000082D05	1300	PA4256	5171	#N/A	#N/A
P1M10000082E05	1301	PA4246	5162	#N/A	#N/A
P1M10000083A11	1302	PA3006	5121	#N/A	#N/A
P1M10000083B01	1303	PA4271	5180	#N/A	#N/A
P1M10000083B12	1304	PA4268	5178	#N/A	#N/A
P1M10000083C11	1305	PA4242	5159	#N/A	#N/A
PIMI0000083C12	1306	PA3006	5121	#N/A	#N/A
P1M10000084A04	1307	PA4942	5201	#N/A	#N/A
P1M10000084D03	1308	PA3006	5121	#N/A	#N/A
P1M10000084E04	1309	PA5493	5218	#N/A	#N/A
P1M10000084E11	1310	PA2196	5102	#N/A	#N/A
P1M10000084F08	1311	PA4271	5180	#N/A	#N/A
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P1M10000086A02	1313	PA4413	5187	#N/A	#N/A
P1M10000086B01	1314	PA4158	5157	#N/A	#N/A
P1M10000086D02	1315	PA2641	5115	#N/A	#N/A
P1M10000086E05	1316	PA3006	5121	#N/A	#N/A
P1M10000087A11	1317	PA4268	5178	#N/A	#N/A
P1M10000087C09	1318	PA2083	5097	#N/A	#N/A
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P1M10000087F09	1321	PA4124	5155	#N/A	#N/A
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P1M10000088A07	1322	PA2742	5120	#N/A	#N/A
P1M10000088D06	1323	PA2108	5099	#N/A	#N/A
P1M10000089C08	1324	PA3048	5125	#N/A	#N/A
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P1M10000089G08	1326	PA2461	5108	#N/A	#N/A
P1M10000090B11	1327	PA3153	5128	#N/A	#N/A
P1M10000090F06	1328	PA2313	5105	#N/A	#N/A
P1M10000090F08	1329	PA4258	5173	#N/A	#N/A
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P1M10000091E09	1330	PA5316	5212	#N/A	#N/A
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Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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P1M10000092D09	1335	PA2128	5100	#N/A	#N/A
P1M10000092E02	1336	PA4256	5171	#N/A	#N/A
P1M10000092F05	1337	PA0423	5067	#N/A	#N/A
P1M10000093A03	1338	PA5088	5205	#N/A	#N/A
P1M10000093B09	1339	PA3703	5138	#N/A	#N/A
P1M10000093C08	1340	PA1868	5092	#N/A	#N/A
P1M10000093E09	1341	PA4332	5183	#N/A	#N/A
P1M10000093F03	1342	PA2101	5098	#N/A	#N/A
P1M10000093H07	1343	PA4665	5195	#N/A	#N/A
P1M10000094F04	1344	PA4268	5178	#N/A	#N/A
P1M10000094H03	1345	PA4744	5198	#N/A	#N/A
P1M10000095C01	1346	PA2488	5110	#N/A	#N/A
P1M10000095C09	1347	PA5443	5216	#N/A	#N/A
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P1M10000095G04	1349	PA4256	5171	#N/A	#N/A
P1M10000096E04	1350	PA0353	5061	#N/A	#N/A
P1M10000096E12	1351	PA4246	5162	#N/A	#N/A
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S1M10000001A05	1354	SAU201508	5819	SAU2c0432 orf 19p	12947
S1M10000001A08	1355	SAU102437	5670	SAU1c0045 orf 33p	12695
S1M10000001A09	1356	SAU101907	5574	SAU1c0040 orf 79p	12442
S1M10000001A10	1357	SAU102602	5708	SAU1c0032_orf_5p	12249
S1M10000001C06	1358	SAU102939	5747	#N/A	#N/A
S1M10000001D01	1359	SAU101907	5574	SAU1c0040_orf_79p	12442
S1M10000001D02	1360	SAU100527	5285	SAU1c0037_orf_101p	12341
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S1M10000001D06	1361	SAU101632	5499	SAU1c0039_orf_3p	12407
S1M10000001D07	1362	SAU101360	5431	SAU1c0044_orf_109p	12555
S1M10000001E02	1363	SAU102602	5708	SAU1c0032_orf_5p	12249
S1M10000001E04	1364	SAU102284	5635	SAU1c0038_orf_5p	12389
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S1M10000001E05	1365	SAU102939	5747	#N/A	#N/A
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S1M10000001E10	1367	SAU103038	5757	#N/A	#N/A
S1M10000001E11	1368	SAU302513	5906	SAU3c1298_orf_lp	13085
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S1M10000001F04	1370	SAU100300	5253	SAU1c0040_orf_90p	12451
S1M10000001F08	1371	SAU102437	5670	SAU1c0045_orf_33p	12695
S1M10000001F09	1372	SAU101907	5574	SAU1c0040_orf_79p	12442
S1M10000001F10	1373	SAU102602	5708	SAU1c0032_orf_5p	12249
S1M10000001F11	1374	SAU102939	5747	#N/A	#N/A
S1M10000001G01	1375	SAU102939	5747	#N/A	#N/A
S1M10000001G07	1376	SAU102939	5747	#N/A	#N/A.
S1M10000001G08	1377	SAU102939	5747	#N/A	#N/A
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S1M10000002A02	1379	SAU102631	5721	SAU1c0045_orf_94p	12712

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M10000002A10	1381	SAU201810	5836	SAU2c0308_orf_2p	12769
S1M10000002A10	1381	SAU202174	5845	SAU2c0412_orf_3p	12895
S1M10000002A10	1381	SAU301148	5888	#N/A	#N/A
S1M10000002A12	1382	SAU200916	5797	SAU2c0373 orf 4p	12838
S1M10000002A12	1382	SAU300455	5872	#N/A	#N/A
S1M10000002A12	1382	SAU301620	5899	SAU3c1478 orf 2p	13140
S1M10000002B01	1383	SAU102602	5708	SAU1c0032 orf 5p	12249
S1M10000002B03	1384	SAU101034	5371	SAU1c0044 orf 27p	12608
S1M10000002B04	1385	SAU101907	5574	SAU1c0040 orf 79p	12442
S1M10000002B05	1386	SAU101868	5565	SAU1c0036 orf 23p	12320
S1M10000002B06	1387	SAU100157	5237	SAU1c0040 orf 81p	12444
S1M10000002B07	1388	SAU101389	5441	SAU1c0038 orf 54p	12387
S1M10000002B09	1389	SAU201810	5836	SAU2c0308 orf 2p	12769
S1M10000002B09	1389	SAU202174	5845	SAU2c0412 orf 3p	12895
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S1M10000002C11	1394	SAU202267	5848	SAU2c0204 orf 2p	12727
S1M10000002C11	1394	SAU202781	5853	SAU2c0109 orf 2p	12718
S1M10000002C11	1394	SAU203001	5859	SAU2c0412 orf 15p	12894
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S1M10000002D02	1397	SAU100741	5318	SAU1c0039 orf 48p	12409
S1M10000002D03	1398	SAU102631	5721	SAU1c0045 orf 94p	12712
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S1M10000002E09	1407	SAU100158	5238	SAU1c0040 orf 80p	12443
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S1M1000002F01	1410	SAU200928	5798	SAU2c0365 orf 5p	12815
S1M1000002F02	1411	SAU301620	5899	SAU3c1478_orf_2p	13140
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TABLE IA

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S1M10000002G05	1417	SAU101907	5574	SAU1c0040_orf_79p	12442
S1M10000002G06	1418	SAU101907	5574	SAU1c0040 orf 79p	12442
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S1M10000002G08	1420	SAU100158	5238	SAU1c0040 orf 80p	12443
S1M10000002G09	1421	SAU102939	5747	#N/A	#N/A
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S1M10000002G11	1423	SAU102939	5747	#N/A	#N/A
S1M10000002G12	1424	SAU101907	5574	SAU1c0040 orf 79p	12442
S1M10000003A01	1425	SAU201810	5836	SAU2c0308 orf 2p	12769
S1M10000003A01	1425	SAU202174	5845	SAU2c0412 orf 3p	12895
S1M10000003A01	1425	SAU301148	5888	#N/A	#N/A
S1M10000003A02	1426	SAU101624	5497	SAU1c0040 orf 25p	12429
S1M10000003A03	1427	SAU101752	5522	SAU1c0040 orf 85p	12447
S1M10000003A04	1428	SAU101360	5431	SAU1c0044 orf 109p	12555
S1M10000003A06	1429	SAU101266	5408	SAU1c0042_orf_117p	12490
S1M10000003A07	1430	SAU101907	5574	SAU1c0040 orf 79p	12442
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S1M10000003A10	1432	SAU100432	5271	SAU1c0040 orf 88p	12450
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S1M10000003B08	1435	SAU100952	5358	SAU1c0043 orf 182p	12523
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S1M10000003B12	1437	SAU302060	5905	SAU3c0879 orf 1p	13042
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S1M10000003C07	1439	SAU101271	5411	SAU1c0037 orf 90p	12366
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S1M10000003E07	1446	SAU100964	5363	SAU1c0044 orf 86p	12641
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S1M10000003F08	1454	SAU102939	5747	#N/A	#N/A
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S1M10000003G04	1457	SAU201810 SAU202174	5845	SAU2c0308_orf_2p	12769
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S1M10000004A11	1463	SAU100521	5283	SAU1c0044_orf_250p	12600
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Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M10000004F07	1497	SAU102764	5734	SAU1c0044_orf_56p	12625
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S1M10000005D07	1535	SAU101869	5566	SAU1c0036_orf_24p	12321
S1M10000005D08	1536	SAU101624	5497	SAUIc0040_orf_25p	12429
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\$1M1000006F04	1584	SAU100964	5363	SAU1c0044_orf_86p	12641
S1M10000006F06	1585	SAU101907	5574	SAU1c0040_orf_79p	12442
S1M1000006G02	1586	SAU101833	5555	SAU1c0038_orf_34p	12373
S1M1000006G03	1587	SAU101400	5444	SAU1c0036_orf_35p	12326
S1M1000006G05	1588	SAU100275	5252	SAU1c0036_orf_15p	12314

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M10000006G09	1591	SAU102939	5747	#N/A	#N/A
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S1M10000006G11	1593	SAU101438	5450	SAU1c0038 orf 40p	12379
S1M10000007A02	1594	SAU102939	5747	#N/A	#N/A
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\$1M10000007B02	1596	SAU202872	5854	SAU2c0393 orf 6p	12866
S1M10000007B11	1597	SAU101476	5459	SAU1c0032 orf 69p	12254
S1M10000007C02	1598	SAU102939	5747	#N/A	#N/A
S1M10000007C04	1599	SAU100608	5297	SAU1c0034 orf 69p	12293
S1M10000007C05	1600	SAU100158	5238	SAU1c0040 orf 80p	12443
\$1M1000007C06	1601	SAU101652	5503	SAU1c0042 orf 123p	12492
S1M10000007C07	1602	SAU101266	5408	SAU1c0042_orf_117p	12490
S1M10000007C08	1603	SAU101717	5513	SAU1c0016_orf_16p	12131
S1M1000007C09	1604	SAU102939	5747	#N/A	#N/A
S1M10000007D03	1605	SAU201810	5836	SAU2c0308 orf 2p	12769
S1M10000007D03	1605	SAU202174	5845	SAU2c0412 orf 3p	12895
S1M10000007D03	1605	SAU301148	5888	#N/A	#N/A
S1M1000007D06	1606	SAU100158	5238	SAU1c0040 orf 80p	12443
S1M1000007D08	1607	SAU102939	5747	#N/A	#N/A
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S1M10000007E04	1610	SAU201810	5836	SAU2c0308 orf 2p	12769
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S1M10000007F10	1618	SAU101791	5532	SAU1c0032_orf_12p	12216
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S1M10000007G03	1622	SAU100952	5358	SAU1c0043 orf 182p	12523
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S1M1000007G07	1623	SAU102652	5725	SAU1c0040_orf_79p	12442
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S1M1000008A04	1627	SAU101491 SAU102939	5747	#N/A	#N/A
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S1M10000008A08	1629	SAU102905 SAU301869	5742	SAU1c0033_orf_45p	1
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Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M10000008B03	1632	SAU103144	5761	SAU1c0045 orf 15p	12663
S1M10000008B04	1633	SAU201810	5836	SAU2c0308_orf_2p	12769
S1M10000008B04	1633	SAU202174	5845	SAU2c0412 orf 3p	12895
S1M10000008B04	1633	SAU301148	5888	#N/A	#N/A
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S1M10000008B08	1635	SAU101652	5503	SAU1c0042 orf 123p	12492
S1M10000008B09	1636	SAU102117	5603	SAU1c0027 orf 6p	12181
S1M10000008B10	1637	SAU100608	5297	SAU1c0034 orf 69p	12293
S1M10000008C05	1638	SAU102939	5747	#N/A	#N/A
S1M10000008C06	1639	SAU102939	5747	#N/A	#N/A
S1M10000008C07	1640	SAU102939	5747	#N/A	#N/A
S1M10000008C08	1641	SAU201571	5824	SAU2c0447_orf_17p	12997
S1M10000008C09	1642	SAU101793	5534	SAU1c0032_orf_14p	12218
S1M10000008D05	1643	SAU100414	5270	SAU1c0022_orf_24p	12148
S1M10000008D09	1644	SAU103038	5757	#N/A	#N/A
S1M10000008E05	1645	SAU101545	5474	SAU1c0037 orf 132p	12348
S1M10000008E08	1646	SAU101907	5574	SAU1c0040 orf 79p	12442
S1M10000008E09	1647	SAU101343	5425	SAU1c0044 orf 40p	12619
S1M10000008E10	1648	SAU101360	5431	SAU1c0044 orf 109p	12555
S1M10000008F01	1649	SAU102284	5635	SAU1c0038 orf 5p	12389
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S1M10000008F02	1650	SAU102007	5590	SAU1c0040 orf 108p	12428
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S1M10000008F06	1652	SAU100741	5318	SAU1c0039 orf 48p	12409
S1M10000008F08	1653	SAU101365	5432	SAU1c0044 orf 112p	12556
S1M10000008F09	1654	SAU201810	5836	SAU2c0308 orf 2p	12769
S1M10000008F09	1654	SAU202174	5845	SAU2c0412 orf 3p	12895
S1M10000008F09	1654	SAU301148	5888	#N/A	#N/A
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S1M1000008G02	1657	SAU201167	5803	SAU2c0407 orf 5p	12887
S1M10000008G03	1658	SAU101637	5500	SAU1c0029 orf 8p	12201
S1M10000008G05	1659	SAU102870	5738	SAU1c0026 orf 17p	12170
S1M10000009A02	1660	SAU101159	5387	SAU1c0036 orf 46p	12331
S1M10000009A04	1661	SAU102979	5750	SAU1c0043 orf 227p	12536
S1M10000009A07	1662	SAU101371	5435	SAU1c0033_orf_7p	12275
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S1M10000009A09	1664	SAU201571	5824	SAU2c0447 orf 17p	12997
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S1M10000009B01	1667	SAU201506	5818	SAU2c0432 orf 18p	12946
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S1M10000009B02	1669	SAU201506	5818	SAU2c0432 orf 18p	12946
S1M1000009B04	1670	SAU102117	5603	SAU1c0027 orf 6p	12181
SIM1000009B05	1671	SAU101752	5522	SAU1c0040 orf 85p	12447
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S1M10000009B10	1674	SAU102527	5693	SAU1c0032_orf_9p	12260
S1M10000009B11	1675	SAU301898	5904	SAU3c1079_orf_lp	13057
S1M10000009B12	1676	SAU102433	5668	SAU1c0045_orf_37p	12701
S1M10000009C01	1677 .	SAU101572	5484	SAU1c0044_orf_211p	12586
S1M10000009C01	1677	SAU101573	5485	SAU1c0044_orf_212p	12587
S1M10000009C02	1678	SAU102418	5664	SAU1c0030_orf_18p	12205
S1M10000009C05	1679	SAU101752	5522	SAU1c0040_orf_85p	12447
S1M10000009C06	1680	SAU102613	5715	SAU1c0041 orf 55p	12475
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S1M10000009C08	1682	SAU100658	5303	SAU1c0038 orf 59p	12388
S1M10000009C09	1683	SAU102129	5604	SAU1c0027 orf 17p	12176
S1M10000009C10	1684	SAU102336	5646	SAU1c0045 orf 146p	12659
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S1M10000009D02	1687	SAU100355	5263	SAU1c0023_orf_6p	12155
S1M10000009D03	1688	SAU102418	5664	SAU1c0030 orf 18p	12205
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S1M10000009D05	1690	SAU100799	5331	SAU1c0045 orf 243p	12682
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S1M10000009D11	1693	SAU200916	5797	SAU2c0373 orf 4p	12838
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SIM1000009F05	1702	SAU101752	5522	SAU1c0025_0ff_18p	12164
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S1M10000009F07	1705	SAU102944	5749	SAU1c0041_orf_51p	12472
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S1M10000009F09	1706	SAU202176 SAU302805	5846	SAU2c0412_orf_3p	13133
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	1707	SAU102392	5658	SAU1c0033_orf_40p	12270
SIM10000009F10	1707	SAU201541	5822	SAU2c0431_orf_14p	12942
S1M10000009G02	1708	SAU101572	5484	SAU1c0044_orf_211p	12586
SIM1000009G02	1708	SAU101573	5485	SAU1c0044_orf_212p	12587
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Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M1000009G09	1713	SAU102693	5731	SAU1c0044_orf_58p	12627
S1M1000009G10	1714	SAU100646	5302	SAU1c0025_orf_5p	12168
S1M1000009G11	1715	SAU100131	5232	SAU1c0043_orf_156p	12517
S1M10000009H01	1716	SAU201506	5818	SAU2c0432_orf_18p	12946
S1M10000009H02	1717	SAU102658	5726	SAU1c0045_orf_121p	12654
S1M10000009H03	1718	SAU201654	5829	SAU2c0442_orf_12p	12982
S1M10000009H05	1719	SAU100582	5292	SAU1c0042 orf 21p	12503
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S1M10000009H05	1719	SAU201929	5838	SAU2c0451 orf 19p	13008
S1M10000009H07	1720	SAU102297	5640	SAU1c0045_orf_41p	12704
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S1M10000011A06	1726	SAU101575	5487	SAU1c0044 orf 214p	12589
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S1M10000011B04	1730	SAU101575	5487	SAU1c0044 orf 214p	12589
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S1M10000011C01	1732	SAU101447	5454	SAU1c0045 orf 244p	12683
S1M10000011C05	1733	SAU100432	5271	SAU1c0040_orf_88p	12450
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S1M10000011D01	1735	SAU101293	5414	SAU1c0044_orf_61p	12631
S1M10000011D02	1736	SAU100414	5270	SAU1c0022_orf_24p	12148
SIM10000011D04	1737	SAU102280	5632	SAU1c0038_orf_3p	12378
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S1M10000011E02	1739	SAU101966	5580	SAU1c0028_orf_41p	12186
S1M10000011E03	1740	SAU101632	5499	SAU1c0039_orf_3p	12407
S1M10000011E04	1741	SAU101572	5484	SAU1c0044_orf_211p	12586
S1M10000011F01	1742	SAU101365	5432	SAU1c0044_orf_112p	12556
S1M10000011F03	1743	SAU102350	5649	SAU1c0040 orf 36p	12433
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S1M10000011G01	1746	SAU301465	5896	SAU3c1429_orf_4p	13121
S1M10000011G03	1747	SAU302626	5907	SAU3c1367_orf_3p	13105
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S1M10000011G05	1749	SAU102350	5649	SAU1c0040_orf_36p	12433
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Сюпе пате	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M10000011H03	1752	SAU202756	5852	SAU2c0470_orf_1p	13027
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S1M10000012A08	1756	SAU101630	5498	SAU1c0039 orf 4p	12410
S1M10000012A08	1756	SAU300156	5867	SAU3c0609 orf 2p	13036
S1M10000012A09	1757	SAU102356	5652	SAU1c0040 orf 41p	12436
SIM10000012A10	1758	SAU101266	5408	SAU1c0042 orf 117p	12490
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S1M10000012A11	1759	SAU200028	5771	SAU2c0145 orf 1p	12721
S1M10000012B01	1760	SAU100751	5321	SAU1c0036 orf 59p	12335
S1M10000012B05	1761	SAU101573	5485	SAU1c0044 orf 212p	12587
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S1M10000012B07	1763	SAU101814	5551	SAU1c0032 orf 32p	12237
S1M10000012B07	1763	SAU101815	5552	SAU1c0032 orf 33p	12238
S1M10000012B11	1764	SAU102551	5698	SAU1c0045 orf 206p	12672
S1M10000012C01	1765	SAU101652	5503	SAU1c0042 orf 123p	12492
S1M10000012C03	1766	SAU100776	5327	SAU1c0041 orf 72p	12482
S1M10000012C04	1767	SAU100776	5327	SAU1c0041 orf 72p	12482
S1M10000012C05	1768	SAU201558	5823	SAU2c0434 orf 5p	12954
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S1M10000012D08	1775	SAU101652	5503	SAU1c0042 orf 123p	12492
SIM10000012D09	1776	SAU101752	5522	SAU1c0040 orf 85p	12447
S1M10000012D12	1777	SAU102620	5718	SAU1c0041 orf 62p	12479
S1M10000012D12	1777	SAU102621	5719	SAU1c0041 orf 63p	12480
S1M10000012D12	1777	SAU202006	5842	SAU2c0456 orf 20p	13018
S1M10000012E01	1778	SAU100733	5314	SAU1c0044 orf 254p	12602
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S1M10000012E02	1779	SAU102485	5686	SAU1c0039 orf 95p	12421
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S1M10000012E07	1781	SAU100390	5267	#N/A	#N/A
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S1M10000012E08	1782	SAU101189	5392	SAU1c0033 orf 25p	12264
S1M10000012E12	1783	SAU201810	5836	SAU2c0308 orf 2p	12769
S1M10000012E12	1783	SAU202174	5845	SAU2c0412_orf_3p	12895
S1M10000012E12	1783	SAU301148	5888	#N/A	#N/A
S1M10000012E12	1784	SAU101793	5534	SAU1c0032 orf 14p	12218
S1M10000012F07	1785	SAU102284	5635	SAU1c0038 orf 5p	12389
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Clone name	Clone SeqID	PathoSeq Locus	Gene SegID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M10000012F09	1787	SAU201403	5815	SAU2c0423 orf 3p	12913
S1M10000012F10	1788	SAU101752	5522	SAU1c0040 orf 85p	12447
S1M10000012F11	1789	SAU101781	5528	SAU1c0037_orf_43p	12353
S1M10000012F12	1790	SAU201810	5836	SAU2c0308 orf 2p	12769
S1M10000012F12	1790	SAU202174	5845	SAU2c0412 orf 3p	12895
S1M10000012F12	1790	SAU301148	5888	#N/A	#N/A
S1M10000012G01	1791	SAU102117	5603	SAU1c0027 orf 6p	12181
S1M10000012G02	1792	SAU301758	5900	SAU3c1508_orf_5p	13156
S1M10000012G03	1793	SAU201301	5809	SAU2c0416_orf_17p	12899
S1M10000012G06	1794	SAU101571	5483	SAU1c0044_orf_210p	12585
S1M10000012G07	1795	SAU101572	5484	SAU1c0044 orf 211p	12586
S1M10000012G07	1795	SAU101573	5485	SAU1c0044 orf 212p	12587
S1M10000012G08	1796	SAU102593	5704	SAU1c0041 orf 39p	12463
S1M10000012G10	1797	SAU100887	5350	SAU1c0018 orf 15p	12138
S1M10000012H05	1798	SAU100157	5237	SAU1c0040 orf 81p	12444
S1M10000012H08	1799	SAU202186	5847	SAU2c0222 orf 1p	12731
S1M10000012H09	1800	SAU100227	5244	SAU1c0043 orf 188p	12525
S1M10000012H10	1801	SAU100432	5271	SAU1c0040 orf 88p	12450
S1M10000012H10	1801	SAU100432	5272	SAU1c0040_orf_88p	12449
S1M10000012H10	1801	SAU101751	5521	SAU1c0040_orf_86p	12448
SIM10000012H11	1802	SAU301118	5886	SAU3c1305 orf 3p	13086
SIM10000013II11	1803	SAU102674	5730	SAU1c0024 orf 12p	12156
SIM10000013A02	1804	SAU101006	5367	SAU1c0024_orf_12p	12136
S1M10000013A05	1805	SAU101000 SAU102450	5675	SAU1c0045 orf 21p	12675
S1M10000013A03	1805	SAU102430 SAU102602	5708	SAU1c0032 orf 5p	12073
S1M10000013A07	1807	SAU102002 SAU101143	5383	SAU1c0032_0ff_3p	12502
S1M10000013A09	1808	SAU101143 SAU101567	5481	SAU1c0022 orf 10p	12302
SIM10000013A09	1808	SAU200030	5772	SAU2c0282 orf 3p	12745
S1M10000013A10	1809	SAU200030 SAU201403	5815	SAU2c0423 orf 3p	12743
SIM10000013A11	1810	SAU101573	5485		12587
SIM10000013A11	1811	SAU101373 SAU100690	5309	SAU1c0044_orf_212p #N/A	#N/A
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				SAU2c0365_orf_5p	12815
S1M10000013B05	1815	SAU100300	5253	SAU1c0040_orf_90p	12451
\$1M10000013B06	1816	SAU100118	5229	SAU1c0015_orf_13p	12125
S1M10000013B07	1817	SAU202174	5845	SAU2c0412_orf_3p	12895
\$1M10000013B07	1817	SAU301148	5888	#N/A	#N/A
S1M10000013B09	1818	SAU200006	5770	SAU2c0157_orf_lp	12723
SIM10000013B11	1819	SAU103042	5758	#N/A	#N/A
S1M10000013C03	1820	SAU101781	5528	SAU1c0037_orf_43p	12353
S1M10000013C05	1821	SAU101038	5372	SAU1c0043_orf_180p	12521
S1M10000013C07	1822	SAU100300	5253	SAU1c0040_orf_90p	12451
S1M10000013C08	1823	SAU101571	5483	SAU1c0044_orf_210p	12585
\$1M10000013C09	1824	SAU102059	5597	SAU1c0034_orf_51p	12286
S1M10000013C10	1825	SAU100736	5316	SAU1c0038_orf_64p	12391
S1M10000013C11	1826	SAU102059	5597	SAU1c0034_orf_51p	12286

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M10000013D08	1828	SAU101798	5538	SAU1c0032_orf_18p	12222
S1M10000013D09	1829	SAU102669	5728	SAU1c0024_orf_7p	12160
S1M10000013D09	1829	SAU302956	5915	SAU3c1513_orf_9p	13161
S1M10000013D11	1830	SAU102433	5668	SAU1c0045 orf 37p	12701
S1M10000013E01	1831	SAU102674	5730	SAU1c0024 orf 12p	12156
S1M10000013E02	1832	SAU101184	5391	SAU1c0035 orf 80p	12305
S1M10000013E04	1833	SAU101802	5542	SAU1c0032 orf 22p	12227
S1M10000013E06	1834	SAU101833	5555	SAU1c0038 orf 34p	12373
S1M10000013E08	1835	SAU100831	5335	SAU1c0038 orf 93p	12403
S1M10000013E09	1836	SAU101571	5483	SAU1c0044 orf 210p	12585
S1M10000013E10	1837	SAU101801	5541	#N/A	#N/A
S1M10000013F02	1838	SAU101570	5482	SAU1c0044 orf 209p	12584
S1M10000013F03	1839	SAU101907	5574	SAU1c0040 orf 79p	12442
S1M10000013F06	1840	SAU103038	5757	#N/A	#N/A
S1M10000013F07	1841	SAU101545	5474	SAU1c0037_orf_132p	12348
S1M10000013F08	1842	SAU100961	5360	SAU1c0044 orf 83p	12638
S1M10000013F09	1843	SAU101398	5442	SAU1c0036 orf 33p	12324
S1M10000013F12	1844	SAU102437	5670	SAU1c0045 orf 33p	12695
S1M10000013G01	1845	SAU100521	5283	SAU1c0044 orf 250p	12600
S1M10000013G04	1846	SAU101592	5490	SAU1c0039 orf 37p	12406
S1M10000013G05	1847	SAU102241	5617	SAU1c0043 orf 25p	12539
S1M10000013G05	1847	SAU102242	5618	SAU1c0043 orf 26p	12540
S1M10000013G06	1848	SAU102380	5654	SAU1c0033 orf 29p	12265
S1M10000013G07	1849	SAU101573	5485	SAU1c0044 orf 212p	12587
S1M10000013G10	1850	SAU201539	5821	SAU2c0431 orf 15p	12943
S1M10000013G11	1851	SAU101890	5570	SAU1c0034 orf 29p	12280
S1M10000013G12	1852	SAU100843	5339	SAU1c0036 orf 40p	12328
S1M10000013H03	1853	SAU100690	5309	#N/A	#N/A
S1M10000013H03	1854	SAU102450	5675	SAU1c0045 orf 21p	12675
S1M10000013H05	1855	SAU200914	5796	SAU2c0373 orf 2p	12837
S1M10000013H07	1856	SAU100414	5270	SAU1c0022 orf 24p	12148
S1M10000013H09	1857	SAU100414	5275	SAU1c0038 orf 67p	12392
S1M10000013H09	1857	SAU200721	5791	SAU2c0339 orf 5p	12797
S1M10000013H10	1858	SAU102059	5597	SAU1c0034 orf 51p	12286
S1M10000013H11	1859	SAU100690	5309	#N/A	#N/A
S1M10000013H11	1860	SAU200564	5784	SAU2c0324 orf 6p	12780
S1M10000014A02	1861	SAU101310	5418	SAU1c0044 orf 125p	12562
S1M10000014A05	1862	SAU101991	5582	SAU1c0044_6H_123p	12362
S1M10000014A07	1863	SAU101526	5470		12434
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	1866			SAU2c0447_orf_17p	12997
S1M10000014B01	1867	SAU100547	5290	SAU1c0032_orf_3p	12240
S1M10000014B02	1868	SAU100432	5271	SAU1c0040_orf_88p	12450
S1M10000014B02	1868	SAU100433	5272	SAU1c0040_orf_87p	12449
S1M10000014B03	1869	SAU100414	5270	SAU1c0022_orf_24p	12148
S1M10000014B04	1870	SAU100778	5328	SAU1c0043_orf_140p	12514

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq
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S1M10000014B06	1872	SAU101199	5395	SAU1c0035_orf_62p	12302
S1M10000014B07	1873	SAU101756	5524	SAU1c0040 orf 82p	12445
S1M10000014B08	1874	SAU101752	5522	SAU1c0040 orf 85p	12447
S1M10000014B10	1875	SAU200006	5770	SAU2c0157 orf 1p	12723
S1M10000014B11	1876	SAU102534	5696	#N/A	#N/A
S1M10000014B12	1877	SAU102534	5696	#N/A	#N/A
S1M10000014C01	1878	SAU101575	5487	SAU1c0044 orf 214p	12589
S1M10000014C05	1879	SAU102602	5708	SAU1c0032 orf 5p	12249
S1M10000014C06	1880	SAU100305	5256	SAU1c0038 orf 77p	12397
S1M10000014C07	1881	SAU101801	5541	#N/A	#N/A
SIM10000014C09	1882	SAU100547	5290	SAU1c0032 orf 3p	12240
S1M10000014C09	1882	SAU102881	5740	SAU1c0032 orf 4p	12242
S1M10000014C10	1883	SAU302901	5912	SAU3c1497 orf 8p	13146
S1M10000014C11	1884	SAU100514	5281	SAU1c0044 orf 57p	12626
S1M10000014C12	1885	SAU101814	5551	SAU1c0032 orf 32p	12237
S1M10000014C12	1885	SAU101815	5552	SAU1c0032 orf 33p	12238
S1M10000014D03	1886	SAU100885	5348	SAU1c0038 orf 38p	12376
S1M10000014D06	1887	SAU100305	5256	SAU1c0038 orf 77p	12397
S1M10000014D08	1888	SAU101752	5522	SAU1c0040 orf 85p	12447
S1M10000014D09	1889	SAU100808	5332	SAU1c0037 orf 12p	12345
S1M10000014D10	1890	SAU102292	5638	SAU1c0038 orf 10p	12368
S1M10000014E01	1891	SAU101793	5534	SAU1c0032 orf 14p	12218
S1M10000014E01	1891	SAU101794	5535	#N/A	#N/A
S1M10000014E04	1892	SAU100964	5363	SAU1c0044 orf 86p	12641
S1M10000014E05	1893	SAU101565	5480	SAU1c0022 orf 8p	12151
S1M10000014E07	1894	SAU100658	5303	SAU1c0038 orf 59p	12388
S1M10000014E07	1894	SAU100659	5304	SAU1c0038 orf 60p	12390
S1M10000014E08	1895	SAU202176	5846	SAU2c0412_orf_3p	12895
S1M10000014E09	1896	SAU102059	5597	SAU1c0034_orf_51p	12286
S1M10000014E09	1896	SAU300269	5869	#N/A	#N/A
S1M10000014E09	1897	SAU102453	5677	SAU1c0045 orf 19p	12669
S1M10000014E12	1898	SAU102284	5635	SAU1c0038 orf 5p	12389
S1M10000014E12	1898	SAU201469	5816	SAU2c0438_orf_6p	12967
S1M10000014F02	1899	SAU100128	5231	#N/A	#N/A
S1M10000014F02	1899	SAU101549	5476	SAU1c0043 orf 64p	12549
S1M10000014F02	1899	SAU101576	5488	SAU1c0044 orf 105p	12543
S1M10000014F03	1900	SAU102200	5611	SAU1c0045_orf_168p	12665
S1M10000014F03	1900	SAU102201	5612	SAU1c0045_orf_169p	12666
S1M10000014F04	1901	SAU102449	5674	SAU1c0045_orf_22p	12677
S1M10000014F04	1901	SAU200914	5796	SAU2c0373 orf 2p	12837
SIM10000014F03	1902	SAU102433	5668	SAU1c0045 orf 37p	12701
S1M10000014F08	1903	SAU102059	5597	SAU1c0045_orf_51p	12701
S1M10000014F09	1904	SAU300269	5869	#N/A	#N/A
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				SAU1c0039_orf_74p	3
S1M10000014G04	1907	SAU101242	5404	SAU1c0044_orf_18p	12578
S1M10000014G06	1908	SAU100275	5252	SAU1c0036_orf_15p	12314

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
S1M10000014G07	1909	SAU201620	5827	#N/A	#N/A
S1M10000014G08	1910	SAU100157	5237	SAU1c0040_orf_81p	12444
S1M10000014G12	1911	SAU102602	5708	SAU1c0032_orf_5p	12249
S1M10000014H02	1912	SAU100242	5246	SAU1c0036_orf_5p	12336
S1M10000014H03	1913	SAU102264	5628	SAU1c0032_orf_60p	12250
S1M10000014H04	1914	SAU100275	5252	SAU1c0036_orf_15p	12314
S1M10000014H05	1915	SAU102116	5602	SAU1c0027_orf_5p	12180
S1M10000014H06	1916	SAU100275	5252	SAU1c0036_orf_15p	12314
S1M10000014H07	1917	SAU103038	5757	#N/A	#N/A
S1M10000014H08	1918	SAU100157	5237	SAU1c0040_orf_81p	12444
S1M10000014H11	1919	SAU102534	5696	#N/A	#N/A
S1M10000015A02	1920	SAU100865	5343	SAU1c0044_orf_99p	12648
S1M10000015A03	1921	SAU102388	5655	SAU1c0033_orf_35p	12267
S1M10000015A05	1922	SAU101815	5552	SAU1c0032_orf_33p	12238
S1M10000015A06	1923	SAU101857	5560	SAU1c0044_orf_156p	12569
S1M10000015A09	1924	SAU100414	5270	SAU1c0022 orf 24p	12148
S1M10000015A10	1925	SAU103038	5757	#N/A	#N/A
S1M10000015A11	1926	SAU101791	5532	SAU1c0032 orf 12p	12216
S1M10000015A12	1927	SAU100158	5238	SAU1c0040_orf_80p	12443
S1M10000015B02	1928	SAU102340	5647	SAU1c0045 orf 149p	12660
S1M10000015B05	1929	SAU103038	5757	#N/A	#N/A
S1M10000015B08	1930	SAU101791	5532	SAU1c0032 orf 12p	12216
S1M10000015B08	1930	SAU101792	5533	SAU1c0032 orf 13p	12217
S1M10000015B09	1931	SAU102585	5703	SAU1c0044 orf 289p	12611
S1M10000015B09	1931	SAU201773	5834	SAU2c0446 orf 4p	12996
S1M10000015B09	1931	SAU302685	5908	SAU3c1403 orf 1p	13113
S1M10000015B10	1932	SAU102308	5642	SAU1c0045_orf_50p	12706
S1M10000015C01	1933	SAU100158	5238	SAU1c0040_orf_80p	12443
S1M10000015C02	1934	SAU102340	5647	SAU1c0045_orf_149p	12660
SIM10000015C03	1935	SAU102390	5657	SAU1c0033_orf_38p	12269
S1M10000015C03	1935	SAU201333	5810	SAU2c0418_orf_8p	12905
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S1M10000015C08	1938	SAU100133	5233	SAU1c0044_orf_170p	12574
S1M10000015C08	1938	SAU100323	5261	SAU1c0044_orf_171p	12575
S1M10000015C10	1939	SAU100414	5270	SAU1c0022_orf_24p	12148
S1M10000015C12	1940	SAU100305	5256	SAU1c0038 orf 77p	12397
S1M10000015D02	1941	SAU100794	5330	SAU1c0028 orf 53p	12189
S1M10000015D03	1942	SAU102032	5591	SAU1c0029_orf_47p	12198
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S1M10000015D05	1944	SAU100793	5329	SAU1c0028 orf 52p	12188
S1M10000015D06	1945	SAU100736	5316	SAU1c0038_orf_64p	12391
S1M10000015D12	1946	SAU101814	5351	SAU1c0032_orf_32p	12237
S1M10000015E02	1947	SAU102390	5657	SAU1c0033 orf 38p	12269
S1M10000015E02	1947	SAU201333	5810	SAU2c0418 orf 8p	12905
S1M10000015E03	1948	SAU200468	5781	SAU2c0429 orf 19p	12937
S1M10000015E06	1949	SAU101320	5420	SAU1c0015 orf 16p	12128
S1M10000015E07	1950	SAU101545	5474	SAU1c0037 orf 132p	12348

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
S1M10000015E09	1951	SAU102433	5668	SAU1c0045_orf_37p	12701
S1M10000015E10	1952	SAU100114	5228	SAU1c0043_orf_225p	12535
S1M10000015E11	1953	SAU102286	5636	SAU1c0038_orf_6p	12393
S1M10000015E11	1953	SAU102287	5637	SAU1c0038_orf_7p	12398
S1M10000015E12	1954	SAU102352	5650	SAU1c0040_orf_38p	12434
S1M10000015F01	1955	SAU100123	5230	SAU1c0043_orf_189p	12526
S1M10000015F01	1955	SAU102001	5586	SAU1c0040 orf 102p	12424
S1M10000015F01	1955	SAU103159	5762	SAU1c0045_orf_204p	12670
S1M10000015F01	1955	SAU201827	5837	SAU2c0449_orf_21p	13002
S1M10000015F02	1956	SAU101561	5479	SAU1c0022_orf_4p	12149
S1M10000015F03	1957	SAU201403	5815	SAU2c0423 orf 3p	12913
S1M10000015F04	1958	SAU201403	5815	SAU2c0423_orf_3p	12913
S1M10000015F06	1959	SAU201385	5814	#N/A	#N/A
S1M10000015F07	1960	SAU101752	5522	SAU1c0040_orf_85p	12447
S1M10000015F08	1961	SAU102102	5600	SAU1c0045_orf_340p	12696
S1M10000015F09	1962	SAU101800	5540	SAU1c0032_orf_20p	12225
S1M10000015F09	1962	SAU101801	5541	#N/A	#N/A
S1M10000015F10	1963	SAU100114	5228	SAU1c0043_orf_225p	12535
S1M10000015G01	1964	SAU102481	5685	SAU1c0039_orf_99p	12422
S1M10000015G02	1965	SAU200058	5773	SAU2c0134_orf_lp	12719
S1M10000015G02	1965	SAU200059	5774	SAU2c0134_orf_3p	12720
S1M10000015G03	1966	SAU101070	5376	SAU1c0034_orf_60p	12291 .
SIM10000015G04	1967	SAU101242	5404	SAU1c0044_orf_18p	12578
S1M10000015G05	1968	SAU101573	5485	SAU1c0044_orf_212p	12587
S1M10000015G06	1969	SAU101156	5386	SAU1c0036_orf_12p	12311
S1M10000015G07	1970	SAU100158	5238	SAU1c0040_orf_80p	12443
S1M10000015G08	1971	SAU101814	5551	SAU1c0032_orf_32p	12237
S1M10000015G09	1972	SAU102143	5607	SAU1c0041_orf_14p	12458
S1M10000015G09	1972	SAU102144	5608	SAU1c0041_orf_15p	12459
S1M10000015G10	1973	SAU101752	5522	SAU1c0040_orf_85p	12447
S1M10000015G11	1974	SAU100275	5252	SAU1c0036_orf_15p	12314
S1M10000015H04	1975	SAU101801	5541	#N/A	#N/A
S1M10000015H04	1975	SAU101802	5542	SAU1c0032_orf_22p	12227
S1M10000015H06	1976	SAU201385	5814	#N/A	#N/A
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S1M10000016A03	1977	SAU101804	5544	#N/A	#N/A
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\$1M10000016A06	1979	SAU200928	5798	SAU2c0365_orf_5p	12815
\$1M10000016A07	1980	SAU100932	5356	SAU1c0044_orf_308p	12615
S1M10000016A09	1981	SAU101067	5375	SAU1c0034_orf_58p	12290
\$1M10000016A09	1981	SAU300732	5877	SAU3c1116_orf_lp	13061
\$1M10000016A10	1982	SAU101571	5483	SAU1c0044_orf_210p	12585
\$1M10000016A12	1983	SAU100522	5284	SAU1c0044_orf_249p	12599
\$1M10000016B02	1984	SAU102449	5674	SAU1c0045_orf_22p	12677
\$1M10000016B05	1985	SAU101320	5420	SAU1c0015_orf_16p	12128
S1M10000016B06	1986	SAU100432	5271	SAU1c0040_orf_88p	12450
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S1M10000016B08	1988	SAU101491	5464	SAU1c0025_orf_20p	12165
S1M10000016B09	1989	SAU301465	5896	SAU3c1429_orf_4p	13121
S1M10000016B10	1990	SAU101006	5367	SAU1c0028_orf_59p	12190
S1M10000016B11	1991	SAU101242	5404	SAU1c0044_orf_18p	12578
S1M10000016B12	1992	SAU101794	5535	#N/A	#N/A
S1M10000016B12	1992	SAU101795	5536	SAU1c0032_orf_15p	12219
S1M10000016C01	1993	SAU100845	5340	SAU1c0036_orf_41p	12329
S1M10000016C02	1994	SAU102049	5595	SAU1c0039_orf_68p	12416
S1M10000016C04	1995	SAU100921	5355	SAU1c0038_orf_76p	12396
S1M10000016C05	1996	SAU101777	5527	SAU1c0037_orf_39p	12352
S1M10000016C06	1997	SAU201810	5836	SAU2c0308_orf_2p	12769
SIM10000016C06	1997	SAU202174	5845	SAU2c0412_orf_3p	12895
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S1M10000016C08	1998	SAU101491	5464	SAU1c0025_orf_20p	12165
S1M10000016C09	1999	SAU102233	5616	SAU1c0043_orf_20p	12531
S1M10000016C10	2000	SAU201513	5820	SAU2c0432_orf_10p	12944
S1M10000016C10	2000	SAU203196	5861	SAU2c0432 orf 11p	12945
S1M10000016C11	2001	SAU101573	5485	SAU1c0044 orf 212p	12587
SIM10000016C12	2002	SAU101752	5522	SAU1c0040 orf 85p	12447
S1M10000016D01	2003	SAU102355	5651	SAU1c0040 orf 40p	12435
S1M10000016D02	2004	SAU200242	5777	SAU2c0250 orf 2p	12734
S1M10000016D04	2005	SAU100921	5355	SAU1c0038 orf 76p	12396
S1M10000016D05	2006	SAU100770	5324	#N/A	#N/A
\$1M10000016D06	2007	SAU100952	5358	SAU1c0043 orf 182p	12523
S1M10000016D08	2008	SAU101070	5376	SAU1c0034 orf 60p	12291
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S1M10000016D10	2010	SAU201513	5820	SAU2c0432_orf_10p	12944
S1M10000016D10	2010	SAU203196	5861	SAU2c0432_orf_11p	12945
S1M10000016D11	2011	SAU101573	5485	SAU1c0044 orf 212p	12587
S1M10000016E04	2012	SAU101371	5435	SAU1c0033_orf_7p	12275
S1M10000016E05	2013	SAU101320	5420	SAU1c0015 orf 16p	12128
S1M10000016E06	2014	SAU102639	5724	#N/A	#N/A
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S1M10000016E09	2017	SAU102527	5693	SAU1c0032 orf 9p	12260
S1M10000016E10	2018	SAU102983	5751	SAU1c0045 orf 224p	12676
S1M10000016E11	2019	SAU102281	5633	SAU1c0038 orf 4p	12384
S1M10000016E12	2020	SAU201571	5824	SAU2c0447_orf_17p	12997
S1M10000016F02	2021	SAU102113	5601	SAU1c0027_orf_2p	12178
S1M10000016F02	2021	SAU301223	5889	SAU3c1345_orf_3p	13090
S1M10000016F03	2022	SAU101864	5562	SAU1c0044 orf 163p	12572
S1M10000016F05	2023	SAU201168	5804	SAU2c0407 orf 8p	12889
S1M10000016F06	2024	SAU102407	5662	#N/A	#N/A
S1M10000016F08	2025	SAU101491	5464	SAU1c0025 orf 20p	12165
S1M10000016F09	2026	SAU102527	5693	SAU1c0032 orf 9p	12260
S1M10000016F11	2027	SAU102113	5601	SAU1c0027 orf 2p	12178

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M10000016G03	2029	SAU101300	5415	SAU1c0044_orf_113p	12557
S1M10000016G03	2029	SAU101365	5432	SAU1c0044_orf_112p	12556
S1M10000016G04	2030	SAU102450	5675	SAU1c0045_orf_21p	12675
S1M10000016G05	2031	SAU102292	5638	SAU1c0038_orf_10p	12368
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S1M10000016H04	2033	SAU101545	5474	SAUIc0037_orf_132p	12348
S1M10000016H08	2034	SAU101067	5375	SAU1c0034_orf_58p	12290
S1M10000016H08	2034	SAU300732	5877	SAU3c1116_orf_1p	13061
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S1M10000017A02	2036	SAU101866	5564	SAU1c0036_orf_21p	12319
S1M10000017A03	2037	SAU101545	5474	SAU1c0037_orf_132p	12348
S1M10000017A03	2037	SAU101546	5475	SAU1c0037_orf_133p	12349
S1M10000017A04	2038	SAU102292	5638	SAU1c0038_orf_10p	12368
S1M10000017A08	2039	SAU102117	5603	SAU1c0027_orf_6p	12181
S1M10000017A11	2040	SAU102437	5670	SAU1c0045_orf_33p	12695
S1M10000017A12	2041	SAU301357	5893	SAU3c1394 orf 2p	13111
S1M10000017B02	2042	SAU102242	5618	SAU1c0043 orf 26p	12540
S1M10000017B05	2043	SAU302513	5906	SAU3c1298 orf 1p	13085
S1M10000017B07	2044	SAU101806	5546	SAU1c0032 orf 25p	12230
S1M10000017B08	2045	SAU101546	5475	SAU1c0037 orf 133p	12349
S1M10000017B09	2046	SAU200928	5798	SAU2c0365 orf 5p	12815
S1M10000017B10	2047	SAU101754	5523	SAU1c0040 orf 84p	12446
S1M10000017B11	2048	SAU101754	5523	SAU1c0040 orf 84p	12446
S1M10000017B12	2049	SAU201375	5811	SAU2c0426_orf_4p	12926
S1M10000017C01	2050	SAU101224	5397	SAU1c0044_orf_98p	12647
S1M10000017C03	2051	SAU101910	5576	SAU1c0040_orf_76p	12440
S1M10000017C05	2052	SAU200657	5789	#N/A	#N/A
S1M10000017C08	2053	SAU101890	5570	SAU1c0034_orf_29p	12280
S1M10000017C09	2054	SAU101398	5442	SAU1c0036_orf_33p	12324
S1M10000017C10	2055	SAU102614	5716	SAU1c0041_orf_56p	12476
S1M10000017C10	2055	SAU102615	5717	SAU1c0041_orf_57p	12477
S1M10000017C11	2056	SAU101799	5539	SAU1c0032_orf_19p	12223
S1M10000017C11	2056	SAU101800	5540	SAU1c0032_orf_20p	12225
S1M10000017C12	2057	SAU101782	5529	SAU1c0037_orf_44p	12354
S1M10000017C12	2057	SAU200994	5802	SAU2c0428_orf_4p	12935
S1M10000017D03	2058	SAU101752	5522	SAU1c0040_orf_85p	12447
S1M10000017D09	2059	SAU101799	5539	SAU1c0032_orf_19p	12223
S1M10000017D09	2059	SAU101800	5540	SAU1c0032_orf_20p	12225
S1M10000017D10	2060	SAU100633	5301	SAU1c0043 orf_147p	12515
S1M10000017E04	2061	SAU101801	5541	#N/A	#N/A
S1M10000017E05	2062	SAU102334	5645	SAU1c0045 orf 144p	12658
S1M10000017E08	2063	SAU101198	5394	SAU1c0035_orf_61p	12301
S1M10000017E11	2064	SAU102883	5741	SAU1c0045_orf_38p	12702
S1M10000017F01	2065	SAU100157	5237	SAU1c0040_orf_81p	12444
S1M10000017F04	2066	SAU100140	5235	SAU1c0032_orf_7p	12258
\$1M10000017F04	2066	SAU100141	5236	SAU1c0032_orf_8p	12259

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S1M10000017F06	2068	SAU102356	5652	SAU1c0040 orf 41p	12436
S1M10000017F11	2069	SAU101463	5458	SAU1c0045 orf 232p	12679
S1M10000017G02	2070	SAU102433	5668	SAU1c0045 orf 37p	12701
S1M10000017G05	2071	SAU102259	5624	SAU1c0032_orf_55p	12245
S1M10000017G06	2072	SAU200565	5785	SAU2c0324 orf 7p	12781
S1M10000018A03	2073	SAU100139	5234	SAU1c0032 orf 6p	12255
S1M10000018A03	2073	SAU102602	5708	SAU1c0032 orf 5p	12249
S1M10000018A04	2074	SAU102142	5606	SAU1c0041 orf 13p	12457
S1M10000018A05	2075	SAU100886	5349	SAU1c0018 orf 16p	12139
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S1M10000018A06	2076	SAU100970	5365	SAU1c0043 orf 197p	12529
S1M10000018A08	2077	SAU100139	5234	SAU1c0032_orf_6p	12255
S1M10000018A08	2077	SAU102602	5708	SAU1c0032_orf_5p	12249
S1M10000018A09	2078	SAU102022	5606	SAU1c0041 orf 13p	12457
S1M10000018A10	2079	SAU100866	5344	SAU1c0044 orf 100p	12553
S1M10000018A11	2079	SAU100139	5234	SAU1c0032 orf 6p	12255
S1M10000018A11	2080	SAU100139 SAU102602	5708	SAU1c0032_orf_5p	12249
S1M10000018A11	2080	SAU102802 SAU100886	5349		12139
S1M10000018B02	2081	SAU100886 SAU100887	5350	SAU1c0018_orf_16p	12139
				SAU1c0018_orf_15p	
S1M10000018B03	2082	SAU101839	5556	SAU1c0042_orf_12p	12495
S1M10000018B05	2083	SAU100300	5253	SAU1c0040_orf_90p	12451
S1M10000018B09	2084	SAU100836	5336	SAU1c0031_orf_13p	12212
S1M10000018B09	2084	SAU202731	5850	#N/A	#N/A
S1M10000018B10	2085	SAU100401	5268	SAU1c0044_orf_174p	12576
SIM10000018B10	2085	SAU300335	5870	#N/A	#N/A
SIM10000018B11	2086	SAU100658	5303	SAU1c0038_orf_59p	12388
S1M10000018C01	2087	SAU101752	5522	SAU1c0040_orf_85p	12447
S1M10000018C02	2088	SAU102447	5672	SAU1c0045_orf_24p	12685
S1M10000018C03	2089	SAU100778	5328	SAU1c0043_orf_140p	12514
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S1M10000018C05	2091	SAU103038	5757	#N/A	#N/A
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S1M10000018C08	2093	SAU102257	5623	SAU1c0032_orf_53p	12244
S1M10000018C09	2094	SAU101065	5374	SAU1c0034_orf_56p	12289
S1M10000018C09	2094	SAU102068	5599	SAU1c0034_orf_55p	12288
S1M10000018C10	2095	SAU100112	5227	SAU1c0044_orf_70p	12634
S1M10000018C11	2096	SAU102663	5727	SAU1c0024_orf_2p	12158
S1M10000018C12	2097	SAU101948	5579	SAU1c0045_orf_69p	12709
S1M10000018D01	2098	SAU101452	5455	SAU1c0045_orf_247p	12684
S1M10000018D02	2099	SAU102284	5635	SAU1c0038_orf_5p	12389
S1M10000018D02	2099	SAU201469	5816	SAU2c0438_orf_6p	12967
S1M10000018D03	2100	SAU101793	5534	SAU1c0032_orf_14p	12218
S1M10000018D04	2101	SAU101798	5538	SAU1c0032_orf_18p	12222
S1M10000018D09	2102	SAU101067	5375	SAU1c0034_orf_58p	12290
S1M10000018D10	2103	SAU301898	5904	SAU3c1079_orf_lp	13057
S1M10000018D11	2104	SAU101752	5522	SAU1c0040 orf 85p	12447

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2142

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5351

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12277

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S1M10000019B08	2144	SAU102422	5666	SAU1c0030_orf_22p	12207
S1M10000019B08	2144	SAU102423	5667	SAU1c0030_orf_23p	12208
S1M10000019B09	2145	SAU100182	5241	SAU1c0037_orf_82p	12362
S1M10000019B09	2145	SAU100251	5248	SAU1c0037_orf_83p	12363
S1M10000019B10	2146	SAU101570	5482	SAU1c0044_orf_209p	12584
S1M10000019B11	2147	SAU100879	5345	SAU1c0041_orf_82p	12483
S1M10000019B12	2148	SAU101793	5534	SAU1c0032_orf_14p	12218
S1M10000019C01	2149	SAU100414	5270	SAU1c0022_orf_24p	12148
S1M10000019C04	2150	SAU103175	5764	SAU1c0045_orf_269p	12687
S1M10000019C04	2150	SAU301472	5897	SAU3c1431_orf_4p	13124
SIM10000019C05	2151	SAU101756	5524	SAU1c0040_orf_82p	12445
S1M10000019C06	2152	SAU101790	5531	SAU1c0032_orf_11p	12215
S1M10000019C06	2152	SAU101791	5532	SAU1c0032_orf_12p	12216
S1M10000019C07	2153	SAU101400	5444	SAU1c0036_orf_35p	12326
S1M10000019C08	2154	SAU202126	5844	SAU2c0045_orf_1p	12714
S1M10000019C11	2155	SAU100301	5254	SAU1c0040_orf_91p	12452
S1M10000019C12	2156	SAU102117	5603	SAU1c0027_orf_6p	12181
S1M10000019D01	2157	SAU102270	5631	SAU1c0032_orf_65p	12253
S1M10000019D02	2158	SAU101145	5384	SAU1c0035_orf_43p	12299
S1M10000019D04	2159	SAU102292	5638	SAU1c0038_orf_10p	12368
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\$1M10000019D07	2162	SAU301898	5904	SAU3c1079_orf_lp	13057
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S1M10000019E02	2166	SAU101624	5497	SAU1c0040_orf_25p	12429
S1M10000019E07	2167	SAU102352	5650	SAU1c0040_orf_38p	12434
S1M10000019F01	2168	SAU102241	5617	SAU1c0043_orf_25p	12539
S1M10000019F05	2169	SAU101612	5493	SAU1c0044_orf_7p	12637
S1M10000019F05	2169	SAU202945	5857	SAU2c0394_orf_7p	12868
S1M10000019F06	2170	SAU101864	5562	SAU1c0044_orf_163p	12572
S1M10000019F08	2171	SAU101571	5483	SAU1c0044_orf_210p	12585
S1M10000019F09	2172	SAU100414	5270	SAU1c0022_orf_24p	12148
S1M10000019F11	2173	SAU101242	5404	SAU1c0044_orf_18p	12578
S1M10000019G04	2174	SAU101793	5534	SAU1c0032_orf_14p	12218
S1M10000019G07	2175	SAU100522	5284	SAU1c0044_orf_249p	12599
S1M10000019G09	2176	SAU100300	5253	SAU1c0040_orf_90p	12451
S1M10000019G10	2177	SAU101235	5400	SAU1c0044_orf_11p	12561
S1M10000019G10	2177	SAU101236	5401	SAU1c0044_orf_12p	12564
S1M10000019G11	2178	SAU101802	5542	SAU1c0032_orf_22p	12227
S1M10000019H05	2179	SAU101802	5542	SAU1c0032_orf_22p	12227
S1M10000019H05	2179	SAU101803	5543	SAU1c0032_orf_23p	12228
S1M10000019H08	2180	SAU102449	5674	SAU1c0045_orf_22p	12677
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S1M10000020A07	2183	SAU200030	5772	SAU2c0282_orf_3p	12745
S1M10000020A11	2184	SAU102437	5670	SAU1c0045_orf_33p	12695
S1M10000020A12	2185	SAU101907	5574	SAU1c0040_orf_79p	12442
S1M10000020B02	2186	SAU100475	5276	SAU1c0036_orf_61p	12337
S1M10000020B03	2187	SAU100059	5224	SAU1c0045_orf_10p	12652
S1M10000020B05	2188	SAU301133	5887	SAU3c1311 orf 3p	13087
S1M10000020B06	2189	SAU100747	5320	SAU1c0044_orf_235p	12597
S1M10000020B07	2190	SAU102433	5668	SAU1c0045 orf 37p	12701
S1M10000020B09	2191	SAU101371	5435	SAU1c0033 orf 7p	12275
S1M10000020B12	2192	SAU102143	5607	SAU1c0041 orf 14p	12458
S1M10000020C09	2193	SAU101545	5474	SAU1c0037 orf 132p	12348
S1M10000020C10	2194	SAU101799	5539	SAU1c0032 orf 19p	12223
S1M10000020C10	2194	SAU101800	5540	SAU1c0032 orf 20p	12225
S1M10000020C11	2195	SAU101452	5455	SAU1c0045_orf_247p	12684
S1M10000020D03	2196	SAU101752	5522	SAU1c0040 orf 85p	12447
S1M10000020D04	2197	SAU102481	5685	SAU1c0039 orf 99p	12422
S1M10000020D06	2198	SAU102578	5701	SAU1c0039 orf 61p	12411
S1M10000020D07	2199	SAU100198	5243	SAU1c0009 orf 1p	12120
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S1M10000020F06	2212	SAU101652	5503	SAU1c0042 orf 123p	12492
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S1M10000020F00	2212	SAU200731	5793	SAU2c0352 orf 2p	12493
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S1M10000020G11	2223	SAU101592	5490	SAU1c0039_orf_37p	12406
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S1M10000020H04	2227	SAU101791	5532	SAU1c0032_orf_12p	12216	
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S1M10000021C12	2249	SAU101726	5515	SAU1c0016 orf 7p	12134	
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S1M10000021F07	2267	SAU101383	5438	SAU1c0022 orf 20p	12147
S1M10000021F09	2268	SAU102059	5597	SAU1c0034 orf 51p	12286
S1M10000021F09	2268	SAU301465	5896	SAU3c1429 orf 4p	13121
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S1M10000021G08	2272	SAU100714	5312	SAU1c0044 orf 74p	12635
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S1M10000021H07	2275	SAU101806	5546	SAU1c0032 orf 25p	12230
S1M10000021H08	2276	SAU102059	5597	SAU1c0034 orf 51p	12286
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S1M10000022A05	2280	SAU101807	5547	SAU1c0032 orf 26p	12231
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S1M10000022B10	2290	SAU101546	5475	SAU1c0037 orf 133p	12349
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S1M10000022C02	2293	SAU102059	5597	SAU1c0034 orf 51p	12286
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S1M10000022E05	2309	SAU301465	5896	SAU3c1429_orf_4p	13121
S1M10000022E09	2310	SAU101235	5400	SAU1c0044_orf_11p	12561
S1M10000022E09	2310	SAU101236	5401	SAU1c0044_orf_12p	12564
S1M10000022F04	2311	SAU101592	5490	SAU1c0039_orf_37p	12406
S1M10000022F06	2312	SAU101868	5565	SAU1c0036_orf_23p	12320
S1M10000022F07	2313	SAU102117	5603	SAU1c0027_orf_6p	12181
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S1M10000022F11	2315	SAU101592	5490	SAU1c0039_orf_37p	12406
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S1M10000022G07	2318	SAU100414	5270	SAU1c0022_orf_24p	12148
S1M10000022G08	2319	SAU100557	5291	SAU1c0044_orf_132p	12565
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S1M10000023B07	2333	SAU101857	5560	SAU1c0044_orf_156p	12569
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S1M10000023B08	2334	SAU100141	5236	SAU1c0032 orf 8p	12259
S1M10000023B09	2335	SAU101340	5423	SAU1c0038_orf_82p	12400
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S1M10000023C12	2342	SAU100077	5226	SAU1c0043_orf_178p	12520
S1M10000023D01	2343	SAU100964	5363	SAU1c0044_orf_86p	12641
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S1M10000023D08	2347	SAU100887	5350	SAU1c0018 orf 15p	12138
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S1M10000023E04	2352	SAU102059	5597	SAU1c0034_orf_51p	12286
\$1M10000023E07	2353	SAU101543	5473	SAU1c0037 orf 130p	12346
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S1M10000023F12	2361	SAU102352	5650	SAU1c0040 orf 38p	12434
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S1M10000023H09	2372	SAU101340	5423	SAU1c0038 orf 82p	12400
SIM10000023H10	2372	SAU101365	5432	SAU1c0044 orf 112p	12556
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S1M10000024D03	2389	SAU100140	5235	SAU1c0044_off_7p	12033
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S1M10000024E03	2392	SAU101401 SAU102418	5664		#N/A 12205
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S1M10000024F02	2396	SAU101447	5454	SAU1c0045_orf_244p	12683
S1M10000024F03	2397	SAU102992	5752	SAU1c0044_orf_60p	12630
S1M10000024F05	2398	SAU201197	5806	SAU2c0429_orf_2p	12938
\$1M10000024F08	2399	SAU101726	5515	SAU1c0016_orf_7p	12134
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S1M10000024G06	2402	SAU102418	5664	SAU1c0030_orf_18p	12205
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S1M10000024H08	2410	SAU102003	5588	SAU1c0040_orf_104p	12426
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S1M10000025B02	2416	SAU101808	5548	SAU1c0032_orf_27p	12232
S1M10000025B03	2417	SAU101385	5439	SAU1c0038_orf_50p	12385
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S1M10000025B09	2420	SAU200928	5798	SAU2c0365_orf_5p	12815
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S1M10000025C01	2422	SAU102292	5638	SAU1c0038_orf_10p	12368
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S1M10000025D03	2429	SAU101772	5526	SAU1c0037_orf_34p	12351
S1M10000025D04	2430	SAU100970	5365	SAU1c0043_orf_197p	12529
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Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq
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S1M10000025D09	2433	SAU100522	5284	SAU1c0044 orf 249p	12599
S1M10000025D10	2434	SAU102200	5611	SAU1c0045 orf 168p	12665
S1M10000025D10	2434	SAU102201	5612	SAU1c0045 orf 169p	12666
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S1M10000025E04	2436	SAU100389	5266	SAU1c0034 orf 14p	12279
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S1M10000025F03	2439	SAU102297	5640	SAU1c0045 orf 41p	12704
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S1M10000025F08	2441	SAU200685	5790	SAU2c0344 orf 9p	12801
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S1M10000025F10	2443	SAU101571	5483	SAU1c0044 orf 210p	12585
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S1M10000025G04	2445	SAU300617	5874	SAU3c1046 orf 2p	13056
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S1M10000025H10	2451	SAU101907	5574	SAU1c0040 orf 79p	12442
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	1			SAU1c0045_orf_19p	
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S1M10000026B05	2463	SAU101546	5475	SAU1c0037_orf_133p	12349
S1M10000026B06	2464	SAU101570	5482	SAU1c0044_orf_209p	12584
S1M10000026B07	2465	SAU101341	5424	SAU1c0044_orf_38p	12618
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S1M10000026B10	2466	SAU101592	5490	SAU1c0039_orf_37p	12406
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Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M10000026C07	2471	SAU101842	5557	SAU1c0042_orf_9p	12510
S1M10000026C08	2472	\$AU100139	5234	SAU1c0032 orf 6p	12255
S1M10000026C11	2473	SAU200657	5789	#N/A	#N/A
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S1M10000026D06	2477	SAU100139	5234	SAU1c0032 orf 6p	12255
S1M10000026D07	2478	SAU101815	5552	SAU1c0032 orf 33p	12238
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SIM10000026E01	2482	SAU101543	5473	SAU1c0037 orf 130p	12346
S1M10000026E07	2483	SAU102939	5747	#N/A	#N/A
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S1M10000026E10	2485	SAU101869	5566	SAU1c0036 orf 24p	12321
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SIM10000026E12	2487	SAU101791 SAU100964	5363	SAU1c0044 orf 86p	12641
SIM10000026F01	2487	SAU100964 SAU101784	5530	SAU1c0044_orf_86p	12355
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				SAU1c0032_orf_6p	12255
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S1M10000026F07		SAU101869	5566	SAU1c0036_orf_24p	12321
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S1M10000026F11	2497	SAU102939	5747	#N/A	#N/A
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S1M10000026G06	2503	SAU101784	5530	SAU1c0037_orf_46p	12355
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S1M10000026G09	2505	SAU100542	5288	SAU1c0043_orf_210p	12532
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S1M10000026G10	2506	SAU102812	5736	SAU1c0015_orf_15p	12127
S1M10000026G12	2507	SAU101551	5477	SAU1c0043_orf_67p	12550
S1M10000026H01	2508	SAU101652	5503	SAU1c0042_orf_123p	12492
S1M10000026H02	2509	SAU102355	5651	SAU1c0040_orf_40p	12435
S1M10000026H03	2510	SAU101801	5541	#N/A	#N/A
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S1M10000026H04	2511	SAU202174	5845	SAU2c0412_orf_3p	12895

SIMI0000026H05 2512 SAU101907 5574 SAU1c0040_ort_79p 124	Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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SIMI0000027610 2515 SAU102479 5683 SAU10039_orf_101p 1248 SIMI0000027A04 2516 SAU101756 5524 SAU10040_orf_82p 1248 SIMI0000027A05 2517 SAU101805 5545 SAU10032_orf_24p 1248 SIMI0000027A08 2518 SAU101772 5526 SAU10037_orf_34p 1233 SIMI0000027A01 2519 SAU101551 5477 SAU10043_orf_67p 1253 SIMI0000027B04 2520 SAU102339 5747 #N/A #N/A #N/A SIMI0000027B06 2521 SAU1002339 5747 #N/A #N/A #N/A SIMI0000027B06 2521 SAU100275 5525 SAU10040_orf_80p 1248 SIMI0000027B07 2522 SAU100158 5238 SAU10040_orf_80p 1248 SIMI0000027B08 2523 SAU101807 5547 SAU10032_orf_26p 1225 SIMI0000027B09 2524 SAU101807 5547 SAU10032_orf_26p 1225 SIMI0000027B09 2524 SAU101265 5407 #N/A #N/A #N/A SIMI0000027C02 2526 SAU101327 5421 SAU10044_orf_296p 1261 SIMI0000027C04 2527 SAU201236 5808 SAU2c0409_orf_10p 1288 SIMI0000027C05 2528 SAU10217 5503 SAU10007_orf_66p 1251 SIMI0000027C06 2529 SAU100114 5228 SAU10007_orf_66p 1251 SIMI0000027C08 2530 SAU101807 5547 SAU10032_orf_26p 1225 SIMI0000027C09 2531 SAU101807 5547 SAU10032_orf_26p 1225 SIMI0000027C09 2531 SAU101807 5547 SAU10032_orf_26p 1225 SIMI0000027C09 2531 SAU101807 5547 SAU10032_orf_26p 1225 SIMI0000027D02 2532 SAU101807 5547 SAU10032_orf_26p 1225 SIMI0000027D02 2533 SAU101807 5547 SAU10032_orf_26p 1225 SIMI0000027D08 2534 SAU10034 5548 SAU10043_orf_179 1236 SIMI0000027D05 2534 SAU10344 5312 SAU10043_orf_19 1245 SIMI0000027D06 2538 SAU10354 5584 SAU10043_orf_19 1245 SIMI0000027D06 2538 SAU10354 5584 SAU10043_orf_19 1245 SIMI0000027D06 2534 SAU10338 5757 #N/A #N/A #N/A	S1M10000026H09	2514	SAU202174	5845	SAU2c0412 orf 3p	12895
SIMI0000027A04 2516 SAUI01756 5524 SAUI0040_orf_82p 124 SIMI0000027A05 2517 SAUI01805 5534 SAUI00032_orf_24p 122 SIMI0000027A08 2518 SAUI01772 5526 SAUI00037_orf_34p 123 SIMI0000027A01 2519 SAUI01551 5477 SAUI00037_orf_34p 123 SIMI0000027B04 2520 SAUI02939 5747 \$\frac{1}{2}\$\text{WIAA} \$\frac{1}{2}\$\text{VIAA}	S1M10000026H09	2514	SAU301148	5888	#N/A	#N/A
SIMI0000027A05 2517 SAUI01805 5545 SAUI0032_orf_24fp 1225 SIMI0000027A08 2518 SAUI01772 5526 SAUI0032_orf_34p 1235 SIMI0000027A01 2519 SAUI01551 S477 SAUI00043_orf_67p 1255 SIMI0000027B04 2520 SAUI02939 S747 Fiv.	S1M10000026H10	2515	SAU102479	5683	SAU1c0039 orf 101p	12405
SIMI0000027A08 2518 SAU101772 S526 SAU10037_ort_34p 123: SIMI0000027B04 2520 SAU102939 S747 SAU100043_ort_67p 125: SIMI0000027B06 2521 SAU1002939 S747 SAU100043_ort_67p 125: SIMI0000027B06 2521 SAU100275 S252 SAU100056_ort_15p 123: SIMI0000027B06 2521 SAU100158 5238 SAU100040_ort_80p 124: SIMI0000027B07 2522 SAU10158 5238 SAU100040_ort_80p 124: SIMI0000027B09 2524 SAU102059 5547 SAU100034_ort_67p 122: SIMI0000027B09 2524 SAU102059 5597 SAU100034_ort_67p 122: SIMI0000027B09 2524 SAU101265 5407 #N/A #N/A #N/A SIMI0000027B01 2525 SAU101265 5407 #N/A #N/A SIMI0000027C02 2526 SAU101327 5421 SAU100044_ort_296p 126: SIMI0000027C03 2528 SAU102117 5603 SAU100094_ort_10p 128: SIMI0000027C05 2528 SAU102117 5603 SAU100094_ort_10p 128: SIMI0000027C06 2529 SAU10114 5228 SAU100094_ort_125p 125: SIMI0000027C06 2529 SAU10114 5228 SAU100034_ort_225p 125: SIMI0000027C09 2531 SAU101807 5547 SAU100032_ort_26p 122: SIMI0000027C00 2532 SAU101652 5503 SAU100042_ort_124p 124: SIMI0000027D02 2532 SAU101652 5503 SAU100042_ort_124p 124: SIMI0000027D02 2532 SAU101653 5504 SAU100042_ort_124p 124: SIMI0000027D03 2533 SAU100300 5253 SAU100040_ort_90p 124: SIMI0000027D04 2535 SAU202708 5849 SAU20043_ort_70p 125: SIMI0000027D06 2535 SAU202708 5849 SAU20043_ort_74p 126: SIMI0000027D06 2535 SAU202708 5849 SAU20043_ort_74p 126: SIMI0000027D06 2535 SAU202708 5849 SAU20044_ort_74p 126: SIMI0000027D07 2538 SAU100360 5534 SAU100040_ort_90p 124: SIMI0000027D06 2535 SAU200708 5849 SAU200373_ort_14p 129: SIMI0000027D06 2536 SAU200708 5849 SAU20043_ort_74p 126: SIMI0000027D07 2538 SAU100360 5309 MN/A #M/A	S1M10000027A04	2516	SAU101756	5524	SAU1c0040 orf 82p	12445
SIMI0000027A08 2518 SAU101772 5526 SAU10037_crt_34p 123: SIMI0000027B04 2520 SAU102939 5747 SAU1c0043_ort_67p 125: SIMI0000027B06 2521 SAU102939 5747 SAU1c0043_ort_67p 125: SIMI0000027B06 2521 SAU100275 5252 SAU1c0036_ort_15p 123: SIMI0000027B07 2522 SAU100158 5238 SAU1c0040_ort_80p 124- SIMI0000027B07 2522 SAU101058 5238 SAU1c0040_ort_80p 124- SIMI0000027B09 2524 SAU102059 5547 SAU1c0034_ort_51p 123: SIMI0000027B09 2524 SAU102059 5597 SAU1c0034_ort_51p 122: SIMI0000027B09 2524 SAU102059 5597 SAU1c0034_ort_51p 122: SIMI0000027B09 2524 SAU101265 5407 #N/A #N/A #N/A SIMI0000027C02 2526 SAU101327 5421 SAU1c0044_ort_296p 126: SIMI0000027C05 2528 SAU1011327 5421 SAU1c0044_ort_296p 126: SIMI0000027C05 2528 SAU102117 5603 SAU1c0027_ort_62p 121: SIMI0000027C06 2529 SAU100114 52228 SAU1c0034_ort_525p 125: SIMI0000027C06 2529 SAU100114 52228 SAU1c0032_ort_62p 122: SIMI0000027C09 2531 SAU101807 5547 SAU1c0032_ort_62p 122: SIMI0000027C00 2532 SAU101545 5474 SAU1c0032_ort_126p 122: SIMI0000027C02 2532 SAU101652 5503 SAU1c0042_ort_123p 124: SIMI0000027D02 25332 SAU101653 5504 SAU1c0042_ort_123p 124: SIMI0000027D02 25334 SAU10533 5504 SAU1c0042_ort_124p 124: SIMI0000027D05 2534 SAU10300 5253 SAU1c0040_ort_90p 124: SIMI0000027D06 2535 SAU202708 5849 SAU2c0385_ort_1p 128: SIMI0000027D06 2535 SAU202708 5849 SAU2c0385_ort_1p 128: SIMI0000027D06 2535 SAU2002708 5849 SAU2c0385_ort_1p 129: SIMI0000027D06 2535 SAU200708 5584 SAU1c0044_ort_74p 126: SIMI0000027D06 2535 SAU200708 5584 SAU1c0044_ort_74p 126: SIMI0000027D06 2538 SAU100590 5309 #N/A	S1M10000027A05	2517	SAU101805	5545	SAU1c0032 orf 24p	12229
SIMI0000027B04 2520 SAUI02939 3747 #N/A #N/A #N/SIMI0000027B06 2521 SAUI00275 5252 SAUI0036_orf_I5p 123: SIMI0000027B07 2522 SAUI00158 5238 SAUIc0036_orf_I5p 123: SIMI0000027B08 2523 SAUI01807 5547 SAUIc0032_orf_26p 122: SIMI0000027B09 2524 SAUI02059 5537 SAUIc0034_orf_51p 122: SIMI0000027B09 2526 SAUI01265 5407 #N/A #N/SIMI000027C02 2526 SAUI01265 5407 #N/A #N/SIMI0000027C04 2527 SAU201236 5808 SAU2c0409_orf_10p 128: SIMI0000027C04 2527 SAU201236 5808 SAU2c0409_orf_10p 128: SIMI0000027C05 2528 SAUI0117 5603 SAUIc0027_orf_5p 121: SIMI0000027C06 2529 SAUI00114 5228 SAUIc0034_orf_25p 125: SIMI0000027C08 2530 SAUI01807 5547 SAUIc0032_orf_26p 122: SIMI0000027C09 2531 SAUI01807 5547 SAUIc0032_orf_26p 123: SIMI0000027C09 2532 SAUI01845 5474 SAUIc0037_orf_132p 123: SIMI0000027D02 2532 SAUI01653 5504 SAUIc0042_orf_123p 124: SIMI0000027D02 2532 SAUI01653 5504 SAUIc0042_orf_123p 124: SIMI0000027D05 2534 SAUI0534 5478 SAUIc0043_orf_70p 125: SIMI0000027D05 2533 SAUI0554 5478 SAUIc0043_orf_70p 125: SIMI0000027D06 2535 SAUI0554 5478 SAUIc0043_orf_70p 125: SIMI0000027D06 2535 SAUI0554 5478 SAUIc0043_orf_70p 125: SIMI0000027D06 2535 SAUI0554 5478 SAUIc0043_orf_70p 125: SIMI0000027D06 2538 SAUI05283 5634 SAUIc0044_orf_74p 126: SIMI0000027D06 2538 SAUI0288 5849 SAU2c0355_orf_1p 129: SIMI0000027D07 2538 SAUI0288 5634 SAUIc0044_orf_74p 126: SIMI0000027D06 2538 SAUI0288 5634 SAUIc0043_orf_19p 124: SIMI0000027D07 2538 SAUI0288 5634 SAUIc0044_orf_74p 126: SIMI0000027D07 2538 SAUI0288 5634 SAUIc0044_orf_74p 126: SIMI0000027D07 2538 SAUI05096 5584 SAUIc0043_orf_5p 129: SIMI0000027D07 2540 SAUI0569 5584 SAUIc0043_orf_5p 123: SIMI0000027D07 2542 SAUI0569 5547 SAUIc0032_orf_26p 122: SIMI0000027E05 2540 SAUI0588 5757 SAUIc0033_orf_35	S1M10000027A08	2518	SAU101772	5526		12351
SIMI0000027B04 2520 SAUI02939 S747 #N/A #N/SIMI0000027B06 2521 SAUI00275 S252 SAUI0036_orf_15p 123 SIMI0000027B07 2522 SAUI00158 S238 SAUIc0046_orf_80p 124 SIMI0000027B08 2523 SAUI01807 S547 SAUIc0032_orf_26p 122: SIMI0000027B09 2524 SAUI02059 S597 SAUIc0034_orf_51p 122: SIMI0000027B09 2524 SAUI01265 S407 #N/A #N/SIMI0000027C02 2526 SAUI01265 S407 #N/A #N/SIMI000027C02 2526 SAUI01327 S421 SAUIc0034_orf_296p 126: SIMI0000027C04 2527 SAU201236 S808 SAU2c049_orf_10p 128: SIMI0000027C05 2528 SAUI0117 S603 SAUIc0027_orf_6p 121: SIMI0000027C06 2529 SAUI0114 S228 SAUIc0034_orf_225p 125: SIMI0000027C08 2530 SAUI01807 S547 SAUIc0032_orf_26p 122: SIMI0000027C09 2531 SAUI01807 S547 SAUIc0032_orf_123p 123: SIMI0000027C09 2532 SAUI01652 S503 SAUIc0037_orf_132p 123: SIMI0000027C02 2532 SAUI01653 S504 SAUIc0032_orf_123p 124: SIMI0000027D02 2533 SAUI0653 S504 SAUIc0042_orf_123p 124: SIMI0000027D05 2533 SAUI0554 S478 SAUIc0043_orf_70p 125: SIMI0000027D05 2533 SAUI0554 S478 SAUIc0043_orf_70p 125: SIMI0000027D06 2535 SAUI0554 S478 SAUIc0043_orf_10p 125: SIMI0000027D06 2535 SAUI0554 S478 SAUIc0043_orf_10p 125: SIMI0000027D06 2535 SAUI00744 S312 SAUIc0044_orf_70p 125: SIMI0000027D06 2538 SAUI0554 S478 SAUIc0043_orf_10p 125: SIMI0000027D06 2538 SAUI02708 S584 SAUIc0043_orf_10p 125: SIMI0000027D06 2538 SAUI0283 S634 SAUIc0044_orf_71p 126: SIMI0000027D07 2538 SAUI0283 S634 SAUIc0044_orf_71p 126: SIMI0000027D06 2538 SAUI0283 S634 SAUIc0044_orf_71p 126: SIMI0000027D07 2540 SAUI06990 S309 #N/A	\$1M10000027A11	2519	SAU101551	5477	1	12550
SIMI0000027B06 2521 SAU100275 5252 SAU10036_orf_15p 123: SIMI0000027B07 2522 SAU10188 5238 SAU1c0040_orf_80p 124: SIMI0000027B08 2523 SAU101807 5547 SAU1c0032_orf_26p 122: SIMI0000027B09 2524 SAU102039 5597 SAU1c0034_orf_51p 122: SIMI0000027B01 2525 SAU101265 5407 #N/A #N/SIMI000027C02 2526 SAU101327 5421 SAU1c004_orf_296p 126: SIMI0000027C02 2526 SAU101327 5421 SAU1c004_orf_10p 128: SIMI0000027C04 2527 SAU201236 5808 SAU2c0409_orf_10p 128: SIMI0000027C05 2528 SAU102117 5603 SAU1c0027_orf_6p 121: SIMI0000027C06 2529 SAU10114 5228 SAU1c0027_orf_6p 123: SIMI0000027C08 2530 SAU101807 5547 SAU1c0032_orf_25p 125: SIMI0000027C09 2531 SAU101807 5547 SAU1c0032_orf_132p 123: SIMI0000027C09 2531 SAU101857 5444 SAU1c0037_orf_132p 123: SIMI0000027D02 2532 SAU101652 5503 SAU1c0042_orf_123p 124: SIMI0000027D02 2532 SAU101653 5504 SAU1c0042_orf_124p 124: SIMI0000027D02 2533 SAU10633 5504 SAU1c0042_orf_124p 124: SIMI0000027D03 2533 SAU10554 5478 SAU1c0043_orf_70p 125: SIMI0000027D05 2534 SAU10554 5484 SAU1c0043_orf_70p 125: SIMI0000027D05 2534 SAU10554 5484 SAU1c0043_orf_74p 126: SIMI0000027D06 2535 SAU202708 5849 SAU2c0335_orf_1p 128: SIMI0000027D09 2537 SAU203524 5864 SAU2c0335_orf_1p 129: SIMI0000027D09 2537 SAU203524 5864 SAU2c0345_orf_1p 129: SIMI0000027D01 2538 SAU10283 5634 SAU1c004_orf_99p 124: SIMI0000027D01 2538 SAU10283 5634 SAU1c004_orf_99p 124: SIMI0000027D05 2540 SAU200916 5797 SAU2c0375_orf_1p 129: SIMI0000027D07 2542 SAU10360 5309 #N/A #N/A #N/A SIMI0000027E05 2540 SAU200547 5259 SAU3012_orf_3p 122: SIMI0000027E05 2540 SAU200547 5259 SAU302_orf_3p 123: SIMI0000027E05 2542 SAU100347 5259 SAU3c032_orf_3p 123: SIMI0000027E05 2548 SAU100347 5259 SAU3c035_orf_3p 123: SIMI0000027E06 2544 SAU10682 5347 SAU1c00	S1M10000027B04	2520	SAU102939	5747		#N/A
SIMI0000027B07 2522 SAU100158 5238 SAU1c0040_orf_80p 124	S1M10000027B06	2521	,	5252	SAU1c0036 orf 15p	12314
SIMI0000027B08 2523 SAU101807 5547 SAU1c0032_ort_Z6p 122: SIMI0000027B09 2524 SAU102059 5597 SAU1c0034_ort_51p 122: SIMI0000027B11 2525 SAU101265 5407 #N/A #N// SIMI0000027C02 2526 SAU101327 5421 SAU1c0044_ort_296p 126: SIMI0000027C04 2527 SAU201236 5808 SAU2c0409_ort_10p 128: SIMI0000027C05 2528 SAU102117 5603 SAU1c0027_ort_6p 121: SIMI0000027C06 2529 SAU100114 5228 SAU100032_ort_26p 122: SIMI0000027C08 2530 SAU101807 5547 SAU1c0032_ort_26p 122: SIMI0000027C09 2531 SAU101807 5547 SAU1c0032_ort_26p 122: SIMI0000027C09 2531 SAU101545 5474 SAU1c0037_ort_132p 123: SIMI0000027D02 2532 SAU101652 5503 SAU1c0042_ort_123p 124: SIMI0000027D02 2532 SAU101653 5504 SAU1c0042_ort_124p 124: SIMI0000027D03 2533 SAU10300 5253 SAU1c0042_ort_124p 124: SIMI0000027D05 2534 SAU101554 5478 SAU1c0042_ort_124p 124: SIMI0000027D05 2534 SAU10300 5253 SAU1c0040_ort_90p 124: SIMI0000027D06 2535 SAU202708 5849 SAU2c0385_ort_1p 128: SIMI0000027D06 2535 SAU203524 5864 SAU2c0385_ort_1p 126: SIMI0000027D09 2537 SAU203524 5864 SAU2c0435_ort_1p 129: SIMI0000027D01 2538 SAU10283 5634 SAU1c0040_ort_9p 124: SIMI0000027D10 2538 SAU10283 5634 SAU1c0040_ort_9p 124: SIMI0000027D10 2538 SAU10283 5634 SAU1c0040_ort_1p 121: SIMI0000027D05 2540 SAU200316 5797 SAU2c0373_ort_4p 126: SIMI0000027E05 2540 SAU20016 5797 SAU2c0373_ort_4p 126: SIMI0000027E05 2540 SAU301620 5899 SAU3c1478_ort_2p 134: SIMI0000027E06 2541 SAU100547 5290 SAU1c0032_ort_3p 122: SIMI0000027E06 2541 SAU100547 5290 SAU1c0032_ort_3p 122: SIMI0000027E07 2542 SAU101807 5547 SAU1c0032_ort_3p 122: SIMI0000027E08 2543 SAU101551 5477 SAU1c0032_ort_3p 122: SIMI0000027E08 2544 SAU100882 5347 SAU1c0038_ort_35p 123: SIMI0000027F06 2549 SAU100882 5347 SAU1c0038_ort_35p 123: SIMI0000027F06 2549 SAU1008		2522	SAU100158	5238		12443
SIM10000027B09 2524 SAU102059 5597 SAU1c0034_orf_S1p 1225 SIM10000027C02 2526 SAU101265 5407 #N/A #N/SIM10000027C02 2526 SAU101265 5407 #N/A #N/SIM10000027C02 2526 SAU101237 5421 SAU1c0044_orf_296p 1266 SIM10000027C04 2527 SAU201236 5808 SAU2c0409_orf_10p 1288 SIM10000027C05 2528 SAU102117 5603 SAU1c0049_orf_20f_6p 1215 SIM10000027C06 2529 SAU100114 5228 SAU1c0043_orf_225p 1255 SIM10000027C08 2530 SAU10807 5547 SAU1c0032_orf_26p 1223 SIM10000027C09 2531 SAU101807 5547 SAU1c0032_orf_132p 1233 SIM10000027C09 2532 SAU101652 55503 SAU1c0042_orf_123p 1245 SIM10000027D02 2532 SAU101652 55503 SAU1c0042_orf_123p 1245 SIM10000027D02 2532 SAU101653 5554 SAU1c0042_orf_124p 1245 SIM10000027D03 2533 SAU100300 5253 SAU1c0040_orf_90p 1245 SIM10000027D05 2534 SAU101554 5478 SAU1c0043_orf_70p 1255 SIM10000027D06 2535 SAU202708 5849 SAU2c0385_orf_1p 1285 SIM10000027D06 2535 SAU202708 5849 SAU2c0385_orf_1p 1285 SIM10000027D06 2538 SAU100714 5312 SAU1c0040_orf_90p 1245 SIM10000027D06 2538 SAU10283 5634 SAU1c0040_orf_90p 1245 SIM10000027D06 2538 SAU10283 5634 SAU1c0040_orf_90p 1245 SIM10000027D06 2538 SAU10283 5634 SAU1c0040_orf_90p 1245 SIM10000027D10 2538 SAU10283 5634 SAU1c0040_orf_90p 1245 SIM10000027D07 2538 SAU100547 5590 SAU2c0373_orf_4p 1285 SIM10000027E05 2540 SAU301620 5899 SAU3c1478_orf_2p 1314 SIM10000027E05 2540 SAU301620 5899 SAU3c1478_orf_2p 1314 SIM10000027E06 2541 SAU100547 5290 SAU1c0032_orf_3p 1224 SIM10000027E07 2542 SAU10547 5290 SAU1c0032_orf_3p 1224 SIM10000027E08 2543 SAU10547 5547 SAU1c0032_orf_3p 1225 SIM10000027E08 2544 SAU105690 5309 #N/A #N/S SIM10000027F06 2548 SAU10582 5347 SAU1c0038_orf_35p 1235 SIM10000027F06 2548 SAU100882 5347 SAU1c0038_orf_35p 123	S1M10000027B08	2523	SAU101807	5547		12231
SIM10000027B11 2525 SAU101265 5407	S1M10000027B09	2524	SAU102059	5597		12286
\$\frac{\text{SIM10000027C02}{\text{CO2}}\$ 2526 \$\text{SAU101327}{\text{SAU201236}}\$ 5421 \$\text{SAU1c0044_orf_296p}\$ 1266 \$\text{SIM10000027C04}{\text{CO2}}\$ 2528 \$\text{SAU201236}{\text{SAU201236}}\$ 5808 \$\text{SAU2c0409_orf_10p}\$ 1288 \$\text{SIM10000027C06}{\text{SIM10000027C06}}\$ 2528 \$\text{SAU102117}{\text{SO3}}\$ 5603 \$\text{SAU1c0027_orf_6p}\$ 1218 \$\text{SIM10000027C06}\$ 2529 \$\text{SAU100114}{\text{SIM10000027C08}}\$ 2530 \$\text{SAU10014}{\text{SIM10000027C08}}\$ 2530 \$\text{SAU1011807}\$ 5547 \$\text{SAU1c0032_orf_26p}\$ 1223 \$\text{SIM10000027C09}\$ 2531 \$\text{SAU101545}\$ 5474 \$\text{SAU1c0037_orf_132p}\$ 1233 \$\text{SIM10000027D02}\$ 2532 \$\text{SAU101652}\$ 5503 \$\text{SAU1c0037_orf_132p}\$ 1245 \$\text{SIM10000027D02}\$ 2532 \$\text{SAU101653}\$ 5504 \$\text{SAU1c0042_orf_124p}\$ 1245 \$\text{SIM10000027D02}\$ 2533 \$\text{SAU100300}\$ 5253 \$\text{SAU1c0040_orf_90p}\$ 1245 \$\text{SIM10000027D03}\$ 2533 \$\text{SAU100300}\$ 5253 \$\text{SAU1c0040_orf_90p}\$ 1255 \$\text{SIM10000027D05}\$ 2534 \$\text{SAU10554}\$ 5478 \$\text{SAU1c0044_orf_70p}\$ 1255 \$\text{SIM10000027D08}\$ 2535 \$\text{SAU202708}\$ 5849 \$\text{SAU2c0385_orf_1p}\$ 1288 \$\text{SIM10000027D09}\$ 2537 \$\text{SAU203524}\$ 5864 \$\text{SAU1c0044_orf_74p}\$ 1265 \$\text{SIM10000027D10}\$ 2538 \$\text{SAU10283}\$ 5634 \$\text{SAU1c0044_orf_99p}\$ 1245 \$\text{SIM10000027D10}\$ 2539 \$\text{SAU10996}\$ 5584 \$\text{SAU1c0044_orf_99p}\$ 1245 \$\text{SIM10000027E05}\$ 2540 \$\text{SAU200916}\$ 5797 \$\text{SAU2c0373_orf_4p}\$ 1285 \$\text{SIM10000027E06}\$ 2541 \$\text{SAU100960}\$ 5309 \$\text{#N/A}\$ \$\text{MN/A}\$ \$\text{#N/A}\$ \$\text{MN/A}\$ \$	S1M10000027B11	2525	SAU101265	5407		#N/A
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SIM10000027E08 2543 SAU201571 5824 SAU20447_orf_17p 1299 SIM10000027E09 2544 SAU101807 5547 SAU1c0032_orf_26p 1223 SIM10000027E11 2545 SAU101551 5477 SAU1c0043_orf_67p 1253 SIM10000027F01 2546 SAU103038 5757 #N/A #N/A SIM10000027F02 2547 SAU101491 5464 SAU1c0025_orf_20p 1216 SIM10000027F05 2548 SAU100882 5347 SAU1c0038_orf_35p 1237 SIM10000027F06 2549 SAU100690 5309 #N/A #N/A SIM10000027F08 2550 SAU200006 5770 SAU2c0157_orf_1p 1272 SIM10000027F09 2551 SAU100858 5341 SAU1c0038_orf_86p 1240 SIM10000027G03 2552 SAU101756 5524 SAU1c0040_orf_82p 1244	SIM10000027E07	2542	SAU100547	5290	SAU1c0032 orf 3p	12240
SIM10000027E09 2544 SAU101807 5547 SAU1c0032_orf_26p 1223 SIM10000027E11 2545 SAU101551 5477 SAU1c0043_orf_67p 1253 SIM10000027F01 2546 SAU103038 5757 #N/A #N/A SIM10000027F02 2547 SAU101491 5464 SAU1c0025_orf_20p 1216 SIM10000027F05 2548 SAU100882 5347 SAU1c0038_orf_35p 1237 SIM10000027F06 2549 SAU100690 5309 #N/A #N/A SIM10000027F08 2550 SAU200006 5770 SAU2c0157_orf_1p 1272 SIM10000027F09 2551 SAU100858 5341 SAU1c0038_orf_86p 1240 SIM10000027G03 2552 SAU101756 5524 SAU1c0040_orf_82p 1244		1				12997
SIM10000027E11 2545 SAU101551 5477 SAU1c0043_orf_67p 1255 SIM10000027F01 2546 SAU103038 5757 #N/A #N/A SIM10000027F02 2547 SAU101491 5464 SAU1c0025_orf_20p 1216 SIM10000027F05 2548 SAU100882 5347 SAU1c0038_orf_35p 1237 SIM10000027F06 2549 SAU100690 5309 #N/A #N/A SIM10000027F08 2550 SAU200006 5770 SAU2c0157_orf_1p 1272 SIM10000027F09 2551 SAU100858 5341 SAU1c0038_orf_86p 1240 SIM10000027G03 2552 SAU101756 5524 SAU1c0040_orf_82p 1244	S1M10000027E09	2544	SAU101807	5547		12231
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S1M10000027F09 2551 SAU100858 5341 SAU1c0038_orf_86p 1240 S1M10000027G03 2552 SAU101756 5524 SAU1c0040_orf_82p 1244						12723
\$\text{S1M10000027G03} \text{2552} \text{SAU101756} \text{5524} \text{SAU1c0040_orf_82p} \text{1244}	I	1.			·	12401
			1			12445
ISTM10000027G04 2553 USAD101777 5527 ISATHe0037 orf 30n 1225	S1M10000027G04	2553	SAU101777	5527	SAU1c0037 orf 39p	12352
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Clone name	Clone SeqID	PathoSeq Locus	Gene SegID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M10000027G09	2557	SAU101807	5547	SAU1c0032_orf_26p	12231
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S1M10000027G11	2558	SAU102534	5696	#N/A	#N/A
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S1M10000027H04	2560	SAU101777	5527	SAU1c0037_orf_39p	12352
S1M10000027H05	2561	SAU102526	5692	SAU1c0045_orf_299p	12691
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S1M10000027H09	2565	SAU101382	5437	SAU1c0022_orf_19p	12146
S1M10000027H10	2566	SAU100158	5238	SAU1c0040_orf_80p	12443
S1M10000027H11	2567	SAU102533	5695	#N/A	#N/A
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S1M10000028A06	2570	SAU100478	5277	SAU1c0044_orf_265p	12605
S1M10000028A06	2570	SAU100996	5366	SAU1c0044_orf_266p	12606
S1M10000028A08	2571	SAU102054	5596	SAUIc0039 orf 74p	12417
S1M10000028B01	2572	SAU101085	5378	SAU1c0034 orf 42p	12284
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S1M10000028B03	2574	SAU100887	5350	SAU1c0018_orf_15p	12138
S1M10000028B04	2575	SAU102764	5734	SAU1c0044_orf_56p	12625
S1M10000028B05	2576	SAU101869	5566	SAU1c0036_orf_24p	12321
S1M10000028B06	2577	SAU201558	5823	SAU2c0434_orf_5p	12954
S1M10000028B08	2578	SAU100158	5238	SAU1c0040_orf_80p	12443
S1M10000028B09	2579	SAU100158	5238	SAU1c0040_orf_80p	12443
S1M10000028C02	2580	SAU203296	5863	SAU2c0442_orf_18p	12983
S1M10000028C04	2581	SAU101381	5436	SAU1c0022_orf_18p	12145
S1M10000028C05	2582	SAU100313	5259	SAU1c0045_orf_153p	12661
S1M10000028C05	2582	SAU100359	5264	SAU1c0032_orf_35p	12239
S1M10000028C05	2582	SAU200297	5778	SAU2c0274_orf_2p	12739
S1M10000028C06	2583	SAU103226	5768	SAU1c0045_orf_95p	12713
S1M10000028C08	2584	SAU101752	5522	SAU1c0040_orf_85p	12447
S1M10000028D03	2585	SAU301898	5904	SAU3c1079_orf_lp	13057
S1M10000028D04	2586	SAU101381	5436	SAU1c0022_orf_18p	12145
S1M10000028D06	2587	SAU200006	5770	SAU2c0157_orf_lp	12723
S1M10000028D07	2588	SAU101271	5411	SAU1c0037_orf_90p	12366
S1M10000028D08	2589	SAU100858	5341	SAU1c0038_orf_86p	12401
S1M10000028D09	2590	SAU100158	5238	SAU1c0040_orf_80p	12443
S1M10000028E01	2591	SAU100062	5225	SAU1c0035_orf_98p	12309
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S1M10000028E03	2592	SAU100770	5324	#N/A	#N/A
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Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID . (protein)	Genemarked gene	full length ORF Protein Seq
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S1M10000028F04	2596	SAU100301	5254	SAU1c0040 orf 91p	12452
S1M10000028F04	2596	SAU100302	5255	SAU1c0040 orf 92p	12453
S1M10000028F05	2597	SAU100301	5254	SAU1c0040 orf 91p	12452
S1M10000028F05	2597	SAU100302	5255	SAU1c0040 orf 92p	12453
S1M10000028F06	2598	SAU100432	5271	SAU1c0040 orf 88p	12450
S1M10000028F06	2598	SAU202756	5852	SAU2c0470 orf 1p	13027
\$1M10000028F07	2599	SAU101006	5367	SAU1c0028 orf 59p	12190
S1M10000028G01	2600	SAU102554	5699	SAU1c0045 orf 209p	12673
S1M10000028G02	2601	SAU201236	5808	SAU2c0409 orf 10p	12891
S1M10000028G02	2601	SAU300338	5871	#N/A	#N/A .
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S1M10000028G04	2603	SAU200916	5797	SAU2c0373_orf_4p	12838
S1M10000028G04	2603	SAU301620	5899 • •	SAU3c1478_orf_2p	13140
S1M10000028G05	2604	SAU100690	5309	#N/A	#N/A
S1M10000028G06	2605	SAU101865	5563	SAU1c0036 orf 20p	12318
S1M10000028G08	2606	SAU101341	5424	SAU1c0044 orf 38p	12618
S1M10000028G08	2606	SAU301275	5892	SAU3c1365 orf 2p	13103
S1M10000028H03	2607	SAU101815	5552	SAU1c0032 orf 33p	12238
S1M10000028H04	2608	SAU103038	5757	#N/A	#N/A
S1M10000028H05	2609	SAU101869	5566	SAU1c0036 orf 24p	12321
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S1M10000029A04	2611	SAU100489	5278	SAU1c0044_orf_133p	12566
S1M10000029A04	2611	SAU100557	5291	SAU1c0044_orf_132p	12565
S1M10000029A09	2612	SAU101495	5467	SAU1c0037_orf_65p	12360
S1M10000029A10	2613	SAU100414	5270	SAU1c0022_orf_24p	12148
S1M10000029A11	2614	SAU101868	5565	SAU1c0036_orf_23p	12320
S1M10000029A12	2615	SAU100865	5343	SAU1c0044_orf_99p	12648
S1M10000029B02	2616	SAU200928	5798	SAU2c0365_orf_5p	12815
S1M10000029B03	2617	SAU201225	5807	SAU2c0412_orf_5p	12896
S1M10000029B04	2618	SAU201621	5828	SAU2c0437_orf_4p	12966
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S1M10000029B06	2620	SAU201571	5824	SAU2c0447_orf_17p	12997
S1M10000029B08	2621	SAU101360	5431	SAU1c0044_orf_109p	12555
S1M10000029B10	2622	SAU101891	5571	SAU1c0034_orf_30p	12281
S1M10000029C02	2623	SAU101271	5411	SAU1c0037_orf_90p	12366
S1M10000029C03	2624	SAU100690	5309	#N/A	#N/A
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S1M10000029C07	2626	SAU102222	5613	SAU1c0043_orf_12p	12511
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\$1M10000029C12	2629	SAU100859	5342	SAU1c0038_orf_87p	12402
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S1M10000029D05	2631	SAU100887	5350	SAU1c0018_orf_15p	12138
S1M10000029D09	2632	SAU101495 .	5467	SAU1c0037_orf_65p	12360
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TABLE IA

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S1M10000029E10	2637	SAU101271	5411	SAU1c0037_orf_90p	12366
S1M10000029E11	2638	SAU101271	5411	SAU1c0037 orf 90p	12366
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S1M10000029F02	2640	SAU101286	5413	SAU1c0034 orf 67p	12292
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S1M10000029F09	2642	SAU301433	5895	SAU3c1420 orf 2p	13118
S1M10000029F10	2643	SAU102621	5719	SAU1c0041 orf 63p	12480
S1M10000029F11	2644	SAU102883	5741	SAU1c0045 orf 38p	12702
S1M10000029F12	2645	SAU102603	5709	SAU1c0041 orf 48p	12469
S1M10000029F12	2645	SAU102609	5713	SAU1c0041_orf_52p	12473
S1M10000029G01	2646	SAU101752	5522	SAU1c0040_orf 85p	12447
S1M10000029G02	2647	SAU101622	5496	SAU1c0040 orf 27p	12430
S1M10000029G03	2648	SAU201571	5824	SAU2c0447 orf 17p	12997
S1M10000029G05	2649	SAU101156	5386	SAU1c0036 orf 12p	12311
S1M10000029G07	2650	SAU101622	5496	SAU1c0040 orf 27p	12430
S1M10000029G08	2651	SAU101365	5432	SAU1c0044 orf 112p	12556
S1M10000029G12	2652	SAU101270	5410	SAU1c0037 orf 89p	12365
S1M10000029H01	2653	SAU100414	5270	SAU1c0022 orf 24p	12148
S1M10000029H05	2654	SAU102613	5715	SAU1c0041 orf 55p	12475
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S1M10000029H08	2656	SAU101271	5411	SAU1c0037 orf 90p	12366
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S1M10000030A02	2659	SAU101543	5473	SAU1c0037_orf_130p	12346
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S1M10000030B07	2666	SAU101180	5389	SAU1c0045 orf 126p	12656
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	1	1	5898	SAU3c1467_orf_2p	13137
S1M10000030C12	2675	SAU100961	5360	SAU1c0044_orf_83p	12638
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S1M10000030D06	2680	SAU102392	5658	SAU1c0033 orf 40p	12270
S1M10000030D06	2680	SAU201541	5822	SAU2c0431 orf 14p	12942
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S1M10000030E12	2689	SAU100300	5253	SAU1c0040 orf 90p	12451
S1M10000030F01	2690	SAU100731	5313	SAU1c0044 orf 252p	12601
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SIM10000031A10	2711	SAU102242 SAU101791	5532	SAU1c0032 orf 12p	12340
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Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M10000031C11	2720	SAU102935	5745	#N/A	#N/A
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S1M10000031D09	2724	SAU102453	5677	SAU1c0045_orf_19p	12669
S1M10000031E02	2725	SAU101350	5429	SAU1c0042_orf_109p	12487
S1M10000031E03	2726	SAU101267	5409	SAU1c0037 orf 86p	12364
S1M10000031E03	2726	SAU300719	5876	SAU3c1108_orf_3p	13059
S1M10000031E04	2727	SAU101752	5522	SAU1c0040_orf_85p	12447
S1M10000031E07	2728	SAU102449	5674	SAU1c0045 orf 22p	12677
S1M10000031E08	2729	SAU100158	5238	SAU1c0040 orf 80p	12443
S1M10000031E10	2730	SAU102433	5668	SAU1c0045_orf_37p	12701
S1M10000031E12	2731	SAU101400	5444	SAU1c0036 orf_35p	12326
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S1M10000032A05	2753	SAU100275	5252	SAU1c0036_orf_15p	12314
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S1M10000032A07	2755	SAU102059	5597	SAU1c0034_orf_51p	12286
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S1M10000032B08	2761	SAU100175	5240	SAU1c0044 orf 204p	12582
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SIM10000032B12	2763	SAU102117	5603	SAU1c0027_orf_6p	12181
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S1M10000032G04	2795	SAU101509	5469	SAU1c0039 orf 81p	12418
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S1M10000032G08	2797	SAU101907	5574	SAU1c0040_orf_79p	12447
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S1M10000032H07	2801	SAU101798	5538	SAU1c0032 orf 18p	12222
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				SAU3c1420_orf_2p	13118
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S1M10000033D10	2819	SAU100813	5334	SAU1c0036_orf_29p	12322
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S1M10000033H02	2837	SAU101907	5574	SAU1c0040 orf 79p	12442
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S1M10000034F09	2889	SAU101869	5566	SAU1c0036_orf_24p	12321
S1M10000034F10	2890	SAU102350	5649	SAU1c0040_orf_36p	12433
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\$1M10000034G09	2897	SAU201775	5835	SAU2c0446 orf 4p	12996
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S1M10000034H03	2902	SAU101571	5483	SAU1c0044 orf 210p	12585
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S1M10000035B08	2917	SAU103232	5769	SAU1c0045_orf_341p	12697
S1M10000035B11	2918	SAU101756	5524	SAU1c0040 orf 82p	12445
S1M10000035C01	2919	SAU200928	5798	SAU2c0365 orf 5p	12815
S1M10000035C02	2920	SAU101039	5373	SAU1c0043 orf 181p	12522
S1M10000035C04	2921	SAU101039 SAU100114	5228	SAU1c0043_0ff_181p	12535
S1M10000035C06	2922	SAU10114 SAU101497	5468	SAU1c0037 orf 66p	12361
S1M10000035C00	2923	SAU101752	5522	SAU1c0040_orf_85p	12361
SIM10000035D01	2923	SAU101732 SAU100414	5270	SAU1c0040_orf_asp	12148
S1M10000035D01	2924	SAU200928	5798	SAU2c0365 orf 5p	12148
S1M10000035D04	2925	SAU102117	5603		12181
SIM1000035D06	2926	SAU102117 SAU100970	5365	SAU1c0027_orf_6p	12181
21W110000032D03	2921	3AU 1009/0	3363	SAU1c0043_orf_197p	12529

TABLE IA

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M10000035E03	2930	SAU102447	5672	SAU1c0045_orf_24p	12685
S1M10000035E04	2931	SAU103025	5755	SAU1c0029_orf_9p	12202
S1M10000035E08	2932	SAU100690	5309	#N/A	#N/A
S1M10000035E09	2933	SAU101197	5393	SAU1c0035_orf_60p	12300
S1M10000035E12	2934	SAU102117	5603	SAU1c0027_orf_6p	12181
S1M10000035F03	2935	SAU101092	5381	SAU1c0028 orf 9p	12192
S1M10000035F03	2935	SAU202882	5855	SAU2c0381 orf 3p	12848
S1M10000035F04	2936	SAU101784	5530	SAU1c0037 orf 46p	12355
S1M10000035F09	2937	SAU203296	5863	SAU2c0442 orf 18p	12983
S1M10000035F12	2938	SAU101427	5447	SAU1c0042 orf 144p	12500
S1M10000035F12	2938	SAU103204	5767	SAU1c0042 orf 143p	12499
S1M10000035G02	2939	SAU101365	5432	SAU1c0044 orf 112p	12556
S1M10000035G09	2940	SAU203296	5863	SAU2c0442 orf 18p	12983
S1M10000035G11	2941	SAU101344	5426	SAU1c0044 orf 41p	12620
S1M10000035G12	2942	SAU101907	5574	SAU1c0040 orf 79p	12442
S1M10000035H01	2943	SAU100140	5235	SAU1c0032 orf 7p	12258
S1M10000035H07	2944	SAU100313	5259	SAU1c0045 orf 153p	12661
S1M10000035H07	2944	SAU100359	5264	SAU1c0032 orf 35p	12239
S1M10000035H07	2944	SAU200297	5778	SAU2c0274 orf 2p	12739
S1M10000035H08	2945	SAU101772	5526	SAU1c0037 orf 34p	12351
S1M10000035H09	2946	SAU100496	5279	SAU1c0041 orf 83p	12484
S1M10000035H09	2946	SAU301004	5882	SAU3c1255 orf 1p	13079
S1M10000035H10	2947	SAU101756	5524	SAU1c0040 orf 82p	12445
S1M10000035H11	2948	SAU101344	5426	SAU1c0044 orf 41p	12620
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S1M10000036A03	2950	SAU101242	5404	SAU1c0044 orf 18p	12578
S1M10000036A04	2951	SAU200994	5802	SAU2c0428 orf 4p	12935
S1M10000036A05	2952	SAU101810	5549	SAU1c0032 orf 28p	12233
S1M10000036A05	2952	SAU101811	5550	SAU1c0032 orf 29p	12234
S1M10000036A05	2952	SAU300110	5865	SAU3c0533 orf 2p	13031
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S1M1000036A11	2954	SAU102117	5603	SAU1c0027_orf_6p	12181
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S1M10000036B04	2956	SAU101571	5483	SAU1c0044_off_200p	12585
S1M10000036B06	2957	SAU101653	5504	SAU1c0042_orf_124p	12493
S1M10000036B07	2958	SAU100887	5350	SAU1c0018 orf 15p	12138
S1M10000036B08	2959	SAU101653	5504	SAU1c0042 orf 124p	12138
S1M10000036B11	2960	SAU101033	5597	SAU1c0034 orf 51p	12493
S1M10000036B11	2960	SAU102039 SAU101791	5532	SAU1c0034_orf_51p	12286
S1M10000036E01	2961	SAU101791 SAU100242	5246	SAU1c0032_orf_12p	12216
S1M10000036C01	2962	SAU100242 SAU101592	5490		12336
		l		SAU1c0039_orf_37p	
S1M10000036C04	2964	SAU102433	5668	SAU1c0045_orf_37p	12701
S1M10000036C05	2965	SAU100497	5280	SAU1c0018_orf_3p	12140
S1M10000036C06	2966	SAU100158	5238	SAU1c0040_orf_80p	12443
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Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M10000036C09	2968	SAU302685	5908	SAU3c1403 orf 1p	13113
S1M10000036C10	2969	SAU100433	5272	SAU1c0040 orf 87p	12449
S1M10000036C10	2969	SAU101751	5521	SAU1c0040 orf 86p	12448
S1M10000036D02	2970	SAU201197	5806	SAU2c0429 orf 2p	12938
S1M10000036D03	2971	SAU103038	5757	#N/A	#N/A
S1M10000036D06	2972	SAU103024	5754	SAU1c0029 orf 6p	12200
S1M10000036D08	2973	SAU101907	5574	SAU1c0040_orf 79p	12442
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S1M10000036D11	2975	SAU101198	5394	SAU1c0035 orf 61p	12301
S1M10000036D12	2976	SAU102117	5603	SAU1c0027_orf_6p	12181
S1M10000036E06	2977	SAU100432	5271	SAU1c0040 orf 88p	12450
S1M10000036E06	2977	SAU202756	5852	SAU2c0470 orf 1p	13027
S1M10000036E08	2978	SAU101028	5370	SAU1c0043 orf 7p	12552
S1M10000036E11	2979	SAU101343	5425	SAU1c0044 orf 40p	12619
S1M10000036F06	2980	SAU101242	5404	SAU1c0044_orf_18p	12578
S1M10000036F07	2981	SAU200928	5798	SAU2c0365 orf 5p	12815
S1M10000036F08	2982	SAU200914	5796	SAU2c0373_orf_2p	12837
S1M10000036F09	2983	SAU100532	5287	SAU1c0044 orf 198p	12580
S1M10000036F10	2984	SAU101586	5489	SAU1c0044 orf 242p	12598
S1M10000036F11	2985	SAU201506	5818	SAU2c0432 orf 18p	12946
S1M10000036G03	2986	SAU101545	5474	SAU1c0037 orf 132p	12348
S1M10000036G07	2987	SAU102355	5651	SAU1c0040 orf 40p	12435
S1M10000036G08	2988	SAU102336	5646	SAU1c0045 orf 146p	12659
S1M10000036G11	2989	SAU101340	5423	SAU1c0038_orf_82p	12400
S1M10000036H01	2990	SAU101793	5534	SAU1c0032 orf 14p	12218
\$1M10000036H02	2991	SAU102117	5603	SAU1c0027 orf 6p	12181
S1M10000036H03	2992	SAU102909	5743	SAU1c0036 orf 16p	12315
S1M10000036H04	2993	SAU102909	5743	SAU1c0036 orf 16p	12315
S1M10000036H05	2994	SAU101798	5538	SAU1c0032_orf_18p	12222
S1M10000036H06	2995	SAU102292	5638	SAU1c0038 orf 10p	12368
S1M10000036H08	2996	SAU102909	5743	SAU1c0036 orf 16p	12315
S1M10000036H11	2997	SAU101653	5504	SAU1c0042 orf 124p	12493
S1M10000037A02	2998	SAU101652	5503	SAU1c0042_orf_123p	12492
S1M10000037A02	2998	SAU101653	5504	SAU1c0042_orf_124p	12493
S1M10000037A03	2999	SAU100128	5231	#N/A	#N/A
S1M10000037A03	2999	SAU101549	5476	SAU1c0043 orf 64p	12549
\$1M10000037A03	2999	SAU101576	5488	SAU1c0044 orf 105p	12554
S1M10000037A06	3000	SAU100964	5363	SAU1c0044 orf 86p	12641
S1M10000037A08	3001	SAU102669	5728	SAU1c0024 orf 7p	12160
S1M10000037A09	3002	SAU101455	5456	SAU1c0045 orf 250p	12686
S1M10000037A09	3002	SAU200916	5797	SAU2c0373 orf 4p	12838
S1M10000037A11	3003	SAU101436	5449	SAU1c0028 orf 23p	12183
SIM10000037A12	3004	SAU200914	5796	SAU2c0373 orf 2p	12837
S1M10000037B03	3005	SAU101999	5585	SAU1c0040 orf 101p	12423

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S1M10000037B06	3008	SAU101806	5546	SAU1c0032 orf 25p	12230
S1M10000037B06	3008	SAU101807	5547	SAU1c0032 orf 26p	12231
S1M10000037B07	3009	SAU101915	5577	SAU1c0040 orf 72p	12439
S1M10000037B08	3010	SAU101592	5490	SAU1c0039 orf 37p	12406
S1M10000037B10	3011	SAU101346	5427	SAU1c0044_orf_43p	12621
S1M10000037B11	3012	SAU101399	5443	SAU1c0036 orf 34p	12325
S1M10000037B12	3013	SAU102117	5603	SAU1c0027 orf 6p	12181
S1M10000037C05	3014	SAU101482	5461	SAU1c0015 orf 10p	12123
S1M10000037C06	3015	SAU101653	5504	SAU1c0042_orf_124p	12493
S1M10000037C07	3016	SAU101641	5501	SAU1c0029 orf 12p	12193
S1M10000037C08	3017	SAU101752	5522	SAU1c0040 orf 85p	12447
S1M10000037C09	3018	SAU101818	5553	SAU1c0038_orf_20p	12369
S1M10000037C10	3019	SAU101752	5522	SAU1c0040 orf 85p	12447
S1M10000037D04	3020	SAU102283	5634	SAU1c0006 orf Ip	12119
S1M10000037D05	3021	SAU100114	5228	SAU1c0043 orf 225p	12535
S1M10000037D06	3022	SAU101996	5584	SAU1c0040 orf 99p	12456
S1M10000037D09	3023	SAU102246	5619	SAU1c0043_orf_30p	12542
S1M10000037D12	3024	SAU101999	5585	SAU1c0040 orf 101p	12423
S1M10000037E02	3025	SAU102447	5672	SAU1c0045 orf 24p	12685
S1M10000037E02	3025	SAU102448	5673	SAU1c0045 orf 23p	12681
S1M10000037E03	3026	SAU100813	5334	SAU1c0036 orf 29p	12322
S1M10000037E06	3027	SAU100921	5355	SAU1c0038 orf 76p	12396
S1M10000037E08	3028	SAU100139	5234	SAU1c0032 orf 6p	12255
S1M10000037E08	3028	SAU100140	5235	SAU1c0032 orf 7p	12258
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S1M10000037E11	3031	SAU201571	5824	SAU2c0447 orf 17p	12997
S1M10000037E12	3032	SAU102602	5708	SAU1c0032 orf 5p	12249
\$1M10000037F02	3033	SAU100776	5327	SAU1c0041 orf 72p	12482
S1M10000037F03	3034	SAU101339	5422	SAU1c0038 orf 81p	12399
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S1M10000037F05	3036	SAU101807	5547	SAU1c0032 orf 26p	12231
S1M10000037F06	3037	SAU102585	5703	SAU1c0044 orf 289p	12611
S1M10000037F06	3037	SAU201773	5834	SAU2c0446 orf 4p	12996
S1M10000037F07	3038	SAU100793	5329	SAU1c0028 orf 52p	12188
S1M10000037F07	3038	SAU301433	5895	SAU3c1420 orf 2p	13118
S1M10000037F08	3039	SAU203001	5859	SAU2c0412 orf 15p	12894
S1M10000037F08	3039	SAU203007	5860	SAU2c0412_orf_10p	12893
S1M10000037F09	3040	SAU101592	5490	SAU1c0039_orf_37p	12406
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S1M10000037F10	3041	SAU102502	5690	SAU1c0045 orf 273p	12689
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S1M10000037G02	3042	SAU102303 SAU100658	5303	SAU1c0045_orf_2/4p	12388
S1M10000037G02	3043	SAU100658 SAU101344	5426	SAU1c0038_orf_39p	12388
S1M10000037G03	3044	SAU101344 SAU101752	5522	SAU1c0044_orf_41p SAU1c0040_orf_85p	12620
S1M10000037G06 S1M10000037G07	3045	SAU101752 SAU103038		SAU1c0040_ori_85p #N/A	
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S1M10000037H02	3049	SAU102059	5597	SAU1c0034 orf 51p	12286
S1M10000037H03	3050	SAU100114	5228	SAU1c0043 orf 225p	12535
S1M10000037H05	3051	SAU100964	5363	SAU1c0044 orf 86p	12641
S1M10000037H07	3052	SAU101571	5483	SAU1c0044_orf 210p	12585
S1M10000037H08	3053	SAU200928	5798	SAU2c0365 orf 5p	12815
S1M10000037H09	3054	SAU100140	5235	SAU1c0032 orf 7p	12258
S1M10000037H11	3055	SAU100608	5297	SAU1c0034 orf 69p	12293
S1M10000038A04	3056	SAU101275	5412	SAU1c0044_orf 257p	12604
S1M10000038A07	3057	SAU100414	5270	SAU1c0022 orf 24p	12148
S1M10000038A08	3058	SAU102059	5597	SAU1c0034 orf 51p	12286
S1M10000038A09	3059	SAU100307	5257	SAU1c0036 orf 134p	12313
\$1M10000038A11	3060	SAU100547	5290	SAU1c0032 orf 3p	12240
S1M10000038A12	3061	SAU101799	5539	SAU1c0032_orf_19p	12223
S1M10000038B01	3062	SAU101483	5462	SAU1c0015_orf_11p	12124
S1M10000038B03	3063	SAU101360	5431	SAU1c0044 orf 109p	12555
S1M10000038B07	3064	SAU102433	5668	SAU1c0045 orf 37p	12701
S1M10000038B08	3065	SAU100308	5258	SAU1c0036 orf 133p	12312
S1M10000038B09	3066	SAU101652	5503	SAU1c0042 orf 123p	12492
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S1M10000038C08	3071	SAU102132	5605	SAU1c0027_orf_19p	12177
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\$1M10000038D07	3077	SAU101652	5503	SAU1c0042 orf 123p	12492
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S1M10000038D11	3081	SAU101365	5432	SAU1c0044 orf 112p	12556
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S1M10000038D12	3082	SAU100952	5358	SAU1c0043 orf 182p	12523
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S1M10000038E02	3084	SAU101842	5557	SAU1c0042 orf 9p	12510
S1M10000038E03	3085	SAU200928	5798	SAU2c0365 orf 5p	12815
S1M10000038E04	3086	SAU101573	5485	SAU1c0044 orf 212p	12513
S1M10000038E05	3087	SAU101653	5504	SAU1c0042 orf 124p	12493
S1M10000038E06	3088	SAU102231	5614	SAU1c0042_0ff_124p	12527
S1M10000030E06	3088	SATI102222	5615	SATT1-0043 orf 10-	12527

5615

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12530

S1M10000038E06

3088

SAU102232

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S1M10000038E12	3091	SAU100838	5337	SAU1c0031_orf_12p	12211
S1M10000038E12	3091	SAU100839	5338	SAU1c0031_orf_11p	12210
S1M10000038F03	3092	SAU102117	5603	SAU1c0027_orf_6p	12181
S1M10000038F04	3093	SAU100964	5363	SAU1c0044_orf_86p	12641
S1M10000038F04	3093	SAU100965	5364	SAU1c0044_orf_87p	12642
S1M10000038F05	3094	SAU100964	5363	SAU1c0044 orf 86p	12641
S1M10000038F05	3094	SAU100965	5364	SAU1c0044 orf 87p	12642
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S1M10000038F08	3096	SAU101752	5522	SAU1c0040 orf 85p	12447
S1M10000038F09	3097	SAU201666	5830	SAU2c0442 orf 11p	12981
S1M10000038F10	3098	SAU101197	5393	SAU1c0035 orf 60p	12300
S1M10000038F11	3099	SAU100747	5320	SAU1c0044_orf_235p	12597
S1M10000038F12	3100	SAU202039	5843	SAU2c0452 orf 20p	13009
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S1M10000038G03	3102	SAU100158	5238	SAU1c0040 orf 80p	12443
S1M10000038G04	3103	SAU100475	5276	SAU1c0036_orf_61p	12337
S1M10000038G06	3104	SAU101189	5392	SAU1e0033 orf 25p	12264
S1M10000038G08	3105	SAU200928	5798	SAU2c0365 orf 5p	12815
S1M10000038G10	3106	SAU102602	5708	SAU1c0032 orf 5p	12249
S1M10000038G11	3107	SAU100123	5230	SAU1c0043 orf 189p	12526
S1M10000038G11	3107	SAU102001	5586	SAU1c0040 orf 102p	12424
S1M10000038G12	3108	SAU101184	5391	SAU1c0035_orf_80p	12305
S1M10000038H03	3109	SAU101798	5538	SAU1c0032 orf 18p	12222
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S1M10000038H09	3111	SAU102340	5647	SAU1c0045 orf 149p	12660
S1M10000038H11	3112	SAU101452	5455	SAU1c0045 orf 247p	12684
S1M10000039A02	3113	SAU100496	5279	SAU1c0041_orf_83p	12484
S1M10000039A02	3113	SAU301004	5882	SAU3c1255 orf lp	13079
S1M10000039A05	3114	SAU100964	5363	SAU1c0044 orf 86p	12641
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S1M10000039A07	3115	SAU100131	5232	SAU1c0043 orf 156p	12517
S1M10000039A08	3116	SAU100522	5284	SAU1c0044 orf 249p	12599
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S1M10000039A12	3118	SAU301465	5896	SAU3c1429 orf 4p	13121
S1M10000039B02	3119	SAU101455	5456	SAU1c0045 orf 250p	12686
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S1M10000039B12	3123	SAU301118	5886	SAU3c1305 orf 3p	13086
S1M10000039C04	3124	SAU102252	5621	SAU1c0032 orf 48p	12241
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S1M10000039C10	3129	SAU101543	5473	SAU1c0037 orf 130p	12346
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S1M10000039D09	3132	SAU102294	5639	SAU1c0044_orf_288p	12610
S1M10000039D09	3132	SAU301080	5885	SAU3c1287_orf_lp	13083
S1M10000039D10	3133	SAU100323	5261	SAU1c0044_orf_171p	12575
S1M10000039E01	3134	SAU102264	5628	SAU1c0032_orf_60p	12250
S1M10000039E08	3135	SAU100412	5269	SAU1c0029_orf_38p	12197
S1M10000039E09	3136	SAU100056	5223	SAU1c0044_orf_176p	12577
S1M10000039E10	3137	SAU102394	5659	SAU1c0033_orf_41p	12271
S1M10000039E10	3137	SAU301118	5886	SAU3c1305_orf_3p	13086
S1M10000039E11	3138	SAU102473	5680	SAU1c0026_orf_30p	12173
S1M10000039F02	3139	SAU201571	5824	SAU2c0447_orf_17p	12997
S1M10000039F03	3140	SAU102527	5693	SAU1c0032_orf_9p	12260
S1M10000039F05	3141	SAU100118	5229	SAU1c0015_orf_13p	12125
S1M10000039F07	3142	SAU102531	5694	SAU1c0045_orf_186p	12667
S1M10000039F08	3143	SAU100158	5238	SAU1c0040_orf_80p	12443
S1M10000039F09	3144	SAU200157	5776	#N/A	#N/A
S1M10000039F10	3145	SAU100059	5224	SAU1c0045_orf_10p	12652
S1M10000039F12	3146	SAU101565	5480	SAU1c0022_orf_8p	12151
S1M10000039G03	3147	SAU101653	5504	SAU1c0042_orf_124p	12493
S1M10000039G04	3148	SAU102292	5638	SAU1c0038_orf_10p	12368
S1M10000039G07	3149	SAU100952	5358	SAU1c0043_orf_182p	12523
S1M10000039G07	3149	SAU101039	5373	SAU1c0043_orf_181p	12522
S1M10000039G10	3150	SAU101815	5552	SAU1c0032_orf_33p	12238
S1M10000039H02	3151	SAU102585	5703	SAU1c0044_orf_289p	12611
S1M10000039H02	3151	SAU201773	5834	SAU2c0446_orf_4p	12996
S1M10000039H03	3152	SAU100313	5259	SAU1c0045_orf_153p	12661
S1M10000039H03	3152	SAU100359	5264	SAU1c0032_orf_35p	12239
S1M10000039H03	3152	SAU200297	5778	SAU2c0274_orf_2p	12739
S1M10000039H04	3153	SAU101752	5522	SAU1c0040_orf_85p	12447
S1M10000039H06	3154	SAU102283	5634	SAU1c0006_orf_lp	12119
S1M10000039H07	3155	SAU100793	5329	SAU1c0028_orf_52p	12188
S1M10000039H07	3155	SAU301433	5895	SAU3c1420_orf_2p	13118
S1M10000039H08	3156	SAU102440	5671	SAU1c0045_orf_30p	12692
S1M10000040A04	3157	SAU100040	5221	SAU1c0043_orf_217p	12533
S1M10000040A05	3158	SAU102671	5729	SAU1c0024_orf_9p	12161
S1M10000040A07	3159	SAU101028	5370	SAU1c0043_orf_7p	12552
S1M10000040A08	3160	SAU200157	5776	#N/A	#N/A
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S1M10000040A11	3162	SAU101801	5541	#N/A	#N/A
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S1M10000040B03	3164	SAU102102	5600	SAU1c0045_orf_340p	12696
S1M10000040B07	3165	SAU101432	5448	SAU1c0028_orf_27p	12184
S1M10000040B11	3166	SAU101198	5394	SAU1c0035_orf_61p	12301
S1M10000040C03	3167	SAU201971	5841	SAU2c0455_orf_17p	13015
S1M10000040C03	3167	SAU301363	5894	#N/A	#N/A
S1M10000040C04	3168	SAU102551	5698	SAU1c0045_orf_206p	12672
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Clone name	Clone	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq
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S1M10000040C07	3171	SAU100970	5365	SAU1c0043 orf 197p	12529
S1M10000040C08	3172	SAU101197	5393	SAU1c0035_orf_60p	12300
S1M10000040C10	3173	SAU201810	5836	SAU2c0308 orf 2p	12769
S1M10000040C10	3173	SAU202174	5845	SAU2c0412 orf 3p	12895
S1M10000040C10	3173	SAU301148	5888	#N/A	#N/A
S1M10000040C11	3174	SAU101869	5566	SAU1c0036 orf 24p	12321
S1M10000040D01	3175	SAU101806	5546	SAU1c0032 orf 25p	12230
S1M10000040D01	3175	SAU101807	5547	SAU1c0032 orf 26p	12231
S1M10000040D03	3176	SAU102200	5611	\$AU1c0045 orf 168p	12665
S1M10000040D03	3176	SAU102201	5612	SAU1c0045 orf 169p	12666
S1M10000040D08	3177	SAU100633	5301	SAU1c0043 orf 147p	12515
S1M10000040D09	3178	SAU101632	5499	SAU1c0039 orf 3p	12407
S1M10000040D11	3179	SAU101546	5475	SAU1c0037 orf 133p	12349
S1M10000040E01	3180	SAU100916	5353	SAU1c0038 orf 71p	12394
S1M10000040E02	3181	SAU101845	5558	SAU1c0042_orf_7p	12506
S1M10000040E04	3182	SAU101546	5475	SAU1c0037 orf 133p	12349
S1M10000040E05	3183	SAU101632	5499	SAU1c0039 orf 3p	12407
S1M10000040E06	3184	SAU101545	5474	SAU1c0037 orf 132p	12348
S1M10000040E07	3185	SAU101006	5367	SAU1c0028 orf 59p	12190
S1M10000040E09	3186	SAU102605	5710	SAU1c0041 orf 49p	12470
S1M10000040E10	3187	SAU100714	5312	SAU1c0044 orf 74p	12635
S1M10000040E11	3188	SAU101226	5398	SAU1c0035 orf 2p	12298
S1M10000040E12	3189	SAU102503	5691	SAU1c0045 orf 274p	12690
S1M10000040E12	3189	SAU201380	5812	SAU2c0426 orf 11p	12922
S1M10000040E01	3190	SAU101226	5398	SAU1c0035 orf 2p	12298
S1M10000040F02	3191	SAU101614	5494	SAU1c0044 orf 9p	12649
SIM10000040F03	3192	SAU101514	5490	SAU1c0039 orf 37p	12406
S1M10000040F04	3193	SAU100123	5230	SAU1c0043 orf 189p	12526
S1M10000040F04	3193	SAU102001	5586	SAU1c0040_orf_102p	12424
S1M10000040F04	3193	SAU102001 SAU103159	5762	SAU1c0045 orf 204p	12670
S1M10000040F04	3193	SAU201827	5837	SAU2c0449 orf 21p	13002
S1M10000040F04	3193	SAU201627 SAU102232	5(15	SAU1c0043 orf 19p	12530
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S1M1000040F09 S1M10000040F12	3197	SAU101610 SAU101752	5522	SAU1c0044_orf_5p	12629
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S1M10000040G02	3200	SAU301773	5901	SAU3c1509_orf_2p	13157
S1M10000040G04	3201	SAU100414	5270	SAU1c0022_orf_24p	12148
S1M10000040G07	3202	SAU101543	5473	SAU1c0037_orf_130p	12346
S1M10000040G08	3203	SAU101752	5522	SAU1c0040_orf_85p	12447
S1M10000040G12	3204	SAU101421	5446	SAU1c0042_orf_138p	12498
S1M10000040H02	3205	SAU100773	5326	SAU1c0038_orf_39p	12377
S1M10000040H03	3206	SAU100414	5270	SAU1c0022_orf_24p	12148
S1M10000040H04	3207	SAU200914	5796	SAU2c0373_orf_2p	12837
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S1M10000041A03	3211	SAU102054	5596	SAU1c0039_orf_74p	12417
S1M10000041B02	3212	SAU101592	5490	SAU1c0039_orf_37p	12406
S1M10000041B03	3213	SAU101592	5490	SAU1c0039_orf_37p	12406
S1M10000041B05	3214	SAU101798	5538	SAU1c0032_orf_18p	12222
S1M10000041B06	3215	SAU301620	5899	SAU3c1478_orf_2p	13140
S1M10000041B07	3216	SAU101145	5384	SAU1c0035_orf_43p	12299
S1M10000041B12	3217	SAU102725	5733	SAU1c0036_orf_68p	12338
S1M10000041C08	3218	SAU102607	5712	SAU1c0041_orf_51p	12472
S1M10000041C08	3218	SAU102944	5749	SAU1c0041_orf_47p	12468
S1M10000041C10	3219	SAU101784	5530	SAU1c0037_orf_46p	12355
S1M10000041C11	3220	SAU101570	5482	SAU1c0044_orf_209p	12584
SIM10000041D06	3221	SAU101777	5527	SAU1c0037 orf 39p	12352
S1M10000041D07	3222	SAU102639	5724	#N/A	#N/A
S1M10000041D08	3223	SAU200030	5772	SAU2c0282 orf 3p	12745
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S1M10000041D12	3225	SAU102658	5726	SAU1c0045 orf 121p	12654
S1M10000041E03	3226	SAU101573	5485	SAU1c0044 orf 212p	12587
S1M10000041E06	3227	SAU101996	5584	SAU1c0040_orf_99p	12456
S1M10000041E09	3228	SAU201236	5808	SAU2c0409 orf 10p	12891
S1M10000041E12	3229	SAU100952	5358	SAU1c0043 orf 182p	12523
S1M10000041F03	3230	SAU101571	5483	SAU1c0044 orf 210p	12585
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S1M10000041F11	3231	SAU102117	5603	SAU1c0027 orf 6p	12181
S1M10000041F12	3232	SAU102480	5684	SAU1c0039 orf 100p	12404
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S1M10000041G11	3237	SAU101802	5542	SAU1c0032 orf 22p	12227
S1M10000041H01	3238	SAU101198	5394	SAU1c0035_orf_61p	12301
S1M10000041H04	3239	SAU100497	5280	SAU1c0018 orf 3p	12140
S1M10000041H05	3240	SAU100242	5246	SAU1c0036 orf 5p	12336
S1M10000041H07	3241	SAU102486	5687	SAU1c0039_orf_93p	12420
S1M10000041H07	3241	SAU102487	5688	SAU1c0039_orf_92p	12419
S1M10000041H08	3242	SAU301133	5887	SAU3c1311 orf 3p	13087
S1M10000041H09	3243	SAU103169	5763	SAU1c0045 orf 230p	12678
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S1M10000042A04	3244	SAU102433	5668	SAU1c0045 orf 37p	12701
S1M10000042A05	3245	SAU102433 SAU102578	5701	SAU1c0045_ort_5/p	12/01
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S1M10000042A07	3247	SAU100633 SAU101495	5467	SAU1c0043_orf_147p	12313
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			5499	SAU1c0039_orf_3p	12407
S1M10000042B02	3251	SAU202736	5851	SAU2c0426_orf_7p	12927
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Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq
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S1M10000042B09	3256	SAU101802	5542	SAU1c0032 orf 22p	12227
S1M10000042B10	3257	SAU100141	5236	SAU1c0032 orf 8p	12259
S1M10000042B10	3257	SAU102527	5693	SAU1c0032 orf 9p	12260
S1M10000042B11	3258	SAU101815	5552	SAU1c0032 orf 33p	12238
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S1M10000042C02	3260	SAU100617	5300	SAU1c0035 orf 102p	12295
S1M10000042C06	3261	SAU102032	5591	SAU1c0029 orf 47p	12198
S1M10000042C10	3262	SAU101495	5467	SAU1c0037 orf 65p	12360
S1M10000042C10	3263	SAU103037	5756	SAU1c0044_orf_303p	12613
S1M10000042C11	3264	SAU103037 SAU101571	5483	SAU1c0044_orf_303p	12515
S1M10000042D04	3265	SAU101371 SAU101632	5499		12383
S1M10000042D07	3265	SAU203296	5863	SAU1c0039_orf_3p	
				SAU2c0442_orf_18p	12983
S1M10000042D11	3267	SAU102663	5727	SAU1c0024_orf_2p	12158
S1M10000042E03	3268	SAU101495	5467	SAU1c0037_orf_65p	12360
S1M10000042E06	3269	SAU102433	5668	SAU1c0045_orf_37p	12701
S1M10000042E08	3270	SAU103198	5766	#N/A	#N/A
S1M10000042F01	3271	SAU102117	5603	SAU1c0027_orf_6p	12181
S1M10000042F02	3272	SAU101891	5571	SAU1c0034_orf_30p	12281
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S1M10000042F06	3274	SAU100773	5326	SAU1c0038_orf_39p	12377
S1M10000042F08	3275	SAU100162	5239	SAU1c0044_orf_206p	12583
S1M10000042F09	3276	SAU100246	5247	SAU1c0042_orf_130p	12496
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S1M10000042F10	3277	SAU102602	5708	SAU1c0032_orf_5p	12249
S1M10000042F11	3278	SAU101653	5504	SAU1c0042_orf_124p	12493
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S1M10000042G08	3281	SAU101907	5574	SAU1c0040_orf_79p	12442
S1M10000042G09	3282	SAU100158	5238	SAU1c0040_orf_80p	12443
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S1M10000042H05	3284	SAU101491	5464	SAU1c0025_orf_20p	12165
S1M10000042H07	3285	SAU100433	5272	SAU1c0040_orf_87p	12449
S1M10000042H11	3286	SAU101632	5499	SAU1c0039_orf_3p	12407
S1M10000043A02	3287	SAU203001	5859	SAU2c0412_orf_15p	12894
S1M10000043A03	3288	SAU101400	5444	SAU1c0036 orf 35p	12326
S1M10000043A04	3289	SAU200088	5775	SAU2c0159 orf 1p	12724
S1M10000043A06	3290	SAU100077	5226	SAU1c0043 orf 178p	12520
S1M10000043A07	3291	SAU101752	5522	SAU1c0040 orf 85p	12447
S1M10000043A08	3292	SAU101543	5473	SAU1c0037 orf 130p	12346
S1M10000043A10	3293	SAU100865	5343	SAU1c0044 orf 99p	12648
S1M10000043A11	3294	SAU100865	5343	SAU1c0044 orf 99p	12648
S1M10000043A12	3295	SAU100887	5350	SAU1c0018 orf 15p	12138
S1M10000043B01	3296	SAU102059	5597	SAU1c0034 orf 51p	12136
S1M10000043B01	3297	SAU100059	5224	SAU1c0034_011_31p	12652
S1M10000043B07	3298	SAU101033	5578	SAU1c0040_orf_66p	12438

Clone name	Clone SeqID	PathoSeq Locus	Gene SeqID (protein)	Genemarked gene	full length ORF Protein Seq ID
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S1M10000043B08	3299	SAU100359	5264	SAU1c0032_orf 35p	12239
S1M10000043B08	3299	SAU200297	5778	SAU2c0274_orf_2p	12739
S1M10000043B09	3300	SAU100521	5283	SAU1c0044_orf 250p	12600
S1M10000043B10	3301	SAU100436	5273	SAU1c0023_orf_20p	12154
S1M10000043B12	3302	SAU102142	5606	SAU1c0041_orf_13p	12457
S1M10000043C02	3303	SAU101777	5527	SAU1c0037_orf 39p	12352
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S1M10000043C12	3306	SAU102059	5597	SAU1c0034 orf 51p	12286
S1M10000043D01	3307	SAU100866	5344	SAU1c0044 orf 100p	12553
S1M10000043D02	3308	SAU301465	5896	SAU3c1429 orf 4p	13121
S1M10000043D04	3309	SAU200928	5798	SAU2c0365 orf 5p	12815
S1M10000043D10	3310	SAU102631	5721	SAU1c0045 orf 94p	12712
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S1M10000043F01	3320	SAU101797	5537	SAU1c0032 orf 17p	12221
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S1M10000047B06	3504	SAU200006	5770	SAU2c0157 orf 1p	12723
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S1M10000047G02	3549	SAU100141	5236	SAU1c0032 orf 8p	12259
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S1M10000048C07	3586	SAU102452	5676	SAU1c0045_orf_20p	12674
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K1M10000003C01	1055	ECO103101	10315	#N/A	#N/A
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K1M10000045A07	1087	ECO104268	10475	#N/A	#N/A
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S4M10000024H02	3736	ECO103738	#N/A	#N/A	#N/A
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S4M10000035E03	3764	KPN103641	#N/A	KPN1c2761 orf 2p	11705
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S4M10000036F07	3768	STY001853	#N/A	STYc00180 orf 22p	#N/A
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S4M10000015E09	3709	STY004152	#N/A	STYc00207_orf_194p	14003
S4M10000016A02	3710	STY004152	#N/A	STYc00207_orf_194p	14003
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TABLE IA

TABLE IC

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KPN107776	3997	5052
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PathoSeq Gene Locus	Nucleotide SeqID	Protein SeqID
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PA0423	4012	5067
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PA0472	4014	5069
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PA0650	4018	5073
PA0715	4019	5074
PA0788	4020	5075
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PA0938	4023	5078
PA1019	4024	5079
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PA1547	4034	5089
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PA1876	4038	5093
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PathoSeq Gene Locus	Nucleotide SeqID	Protein SeqID
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PathoSeq Gene Locus	Nucleotide SeqID	Protein SeqID
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PathoSeq Gene Locus	Nucleotide SeqID	Protein SeqID
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SAU102222	4558	5613
SAU102231	4559	5614
SAU102232	4560	5615
SAU102233	4561	5616
SAU102241	4562	5617
SAU102242	4563	5618
SAU102246	4564	5619
SAU102247	4565	5620
SAU102252	4566	5621
SAU102256	4567	5622
SAU102257	4568	5623
SAU102259	4569	5624
SAU102260	4570	5625
SAU102261	4571	5626
SAU102262	4572	5627
SAU102264	4573	5628
SAU102265	4574	5629
SAU102268	4575	5630
SAU102270	4576	5631
SAU102280	4577	5632
SAU102281	4578	5633
SAU102283	4579	5634
SAU102284	4580	5635
SAU102286	4581	5636
SAU102287	4582	5637
SAU102292	4583	5638
SAU102294	4584	5639
SAU102297	4585	5640
SAU102298	4586	5641
SAU102308	4587	5642

PathoSeq Gene Locus	Nucleotide SeqID	Protein SeqID
SAU102318	4588	5643
SAU102333	4589	5644
SAU102334	4590	5645
SAU102336	4591	5646
SAU102340	4592	5647
SAU102345	4593	5648
SAU102350	4594	5649
SAU102352	4595	5650
SAU102355	4596	5651
SAU102356	4597	5652
SAU102378	4598	5653
SAU102380	4599	5654
SAU102388	4600	5655
SAU102389	4601	5656
SAU102390	4602	5657
SAU102392	4603	5658
SAU102394	4604	5659
SAU102396	4605	5660
SAU102401	4606	5661
SAU102401 SAU102407	4607	5662
SAU102417	4608	5663
SAU102417 SAU102418	4609	5664
SAU102418 SAU102420	4610	5665
SAU102420 SAU102422	4611	5666
SAU102422 SAU102423	4612	5667
SAU102423 SAU102433	4613	5668
SAU102434	4614	5669
SAU102437	4615	5670
SAU102440	4616	5671
SAU102447	4617	5672
SAU102447	4618	5673
SAU102449	4619	5674
SAU102450	4620	5675
SAU102452	4621	5676
SAU102453	4622	5677
SAU102460	4623	5678
SAU102469	4624	5679
SAU102473	4625	5680
SAU102473	4626	5681
SAU102474	4627	5682
SAU102479	4628	5683
SAU102479 SAU102480	4629	5684
SAU102480 SAU102481	4630	5685
SAU102481 SAU102485	4631	5686
SAU102485 SAU102486	4632	5687
SAU102487	4633 4634	5688
SAU102498		5689
SAU102502	4635 4636	5690
SAU102503	4036	5691

PathoSeq Gene Locus	Nucleotide SeqID	Protein SeqID
SAU102526	4637	5692
SAU102527	4638	5693
SAU102531	4639	5694
SAU102533	4640	5695
SAU102534	4641	5696
SAU102541	4642	5697
SAU102551	4643	5698
SAU102554	4644	5699
SAU102575	4645	5700
SAU102578	4646	5701
SAU102584	4647	5702
SAU102585	4648	5703
SAU102593	4649	5704
SAU102598	4650	5705
SAU102599	4651	5706
SAU102601	4652	5707
SAU102602	4653	5708
SAU102603	4654	5709
SAU102605	4655	5710
SAU102606	4656	5711
SAU102607	4657	5712
SAU102609	4658	5713
SAU102610	4659	5714
SAU102613	4660	5715
SAU102614	4661	5716
SAU102615	4662	5717
SAU102620	4663	5718
SAU102621	4664	5719
SAU102629	4665	5720
SAU102631	4666	5721
SAU102636	4667	5722
SAU102637	4668	5723
SAU102639	4669	5724
SAU102652	4670	5725
SAU102658	4671	5726
SAU102663	4672	5727
SAU102669	4673	5728
SAU102671	4674	5729
SAU102674	4675	5730
SAU102693	4676	5731
SAU102694	4677	5732
SAU102725	4678	5733
SAU102764	4679	5734
SAU102766	4680	5735
SAU102812	4681	5736
SAU102863	4682	5737
SAU102870	4683	5738
SAU102880	4684	5739
SAU102881	4685	5740

PathoSeq Gene Locus	Nucleotide SeqID	Protein SeqID
SAU102883	4686	5741
SAU102905	4687	5742
SAU102909	4688	5743
SAU102933	4689	5744
SAU102935	4690	5745
SAU102936	4691	5746
SAU102939	4692	5747
SAU102942	4693	5748
SAU102944	4694	5749
SAU102979	4695	5750
SAU102983	4696	5751
SAU102992	4697	5752
SAU103010	4698	5753
SAU103024	4699	5754
SAU103025	4700	5755
SAU103037	4701	5756
SAU103038	4702	5757
SAU103042	4703	5758
SAU103077	4704	5759
SAU103115	4705	5760
SAU103144	4706	5761
SAU103159	4707	5762
SAU103169	4708	5763
SAU103175	4709	5764
SAU103191	4710	5765
SAU103198	4711	5766
SAU103204	4712	5767
SAU103226	4713	5768
SAU103232	4714	5769
SAU200006	4715	5770
SAU200028	4716	5771
SAU200030	4717	5772
SAU200058	4718	5773
SAU200059	4719	5774
SAU200088	4720	5775
SAU200157	4721	5776
SAU200242	4722	5777
SAU200297	4723	5778
SAU200345	4724	5779
SAU200392	4725	5780
SAU200468	4726	5781
SAU200558	4727	5782
SAU200561	4728	5783
SAU200564	4729	5784
SAU200565	4730	5785
SAU200593	4731	5786
SAU200601	4732	5787
SAU200628	4733	5788
SAU200657	4734	5789

PathoSeq Gene Locus	Nucleotide SeqID	Protein SeqID
SAU200685	4735	5790
SAU200721	4736	5791
SAU200725	4737	5792
SAU200731	4738	5793
SAU200740	4739	5794
SAU200752	4740	5795
SAU200914	4741	5796
SAU200916	4742	5797
SAU200928	4743	5798
SAU200934	4744	5799
SAU200949	4745	5800
SAU200960	4746	5801
SAU200994	4747	5802
SAU201167	4748	5803
SAU201168	4749	5804
SAU201184	4750	5805
SAU201197	4751	5806
SAU201225	4752	5807
SAU201236	4753	5808
SAU201301	4754	5809
SAU201333	4755	5810
SAU201375	4756	5811
SAU201380	4757	5812
SAU201381	4758	5813
SAU201385	4759	5814
SAU201403	4760	5815
SAU201469	4761	5816
SAU201486	4762	5817
SAU201506	4763	5818
SAU201508	4764	5819
SAU201513	4765	5820
SAU201539	4766	5821
SAU201541	4767	5822
SAU201558	4768	5823
SAU201571	4769	5824
SAU201611	4770	5825
SAU201615	4771	5826
SAU201620	4772	5827
SAU201621	4773	5828
SAU201654	4774	5829
SAU201666	4775	5830
SAU201743	4776	5831
SAU201752	4777	5832
SAU201765	4778	5833
SAU201703 SAU201773	4779	5834
SAU201775	4779	5835
SAU201773	4780	5836
SAU201827	4781	5837
SAU201929	4782	5838
5710201929	4703	3030

PathoSeq Gene Locus	Nucleotide SeqID	Protein SeqID
SAU201952	4784	5839
SAU201961	4785	5840
SAU201971	4786	5841
SAU202006	4787	5842
SAU202039	4788	5843
SAU202126	4789	5844
SAU202174	4790	5845
SAU202174	4791	5846
SAU202186	4792	5847
SAU202267	4792	5848
SAU202708	4794	5849
SAU202731	4795	5850
SAU202731	4796	5851
SAU202756	4797	5852
SAU202781	4798	5853
SAU202781 SAU202872		5854
	4799	
SAU202882	4800	5855
SAU202930	4801	5856
SAU202945	4802	5857
SAU202968	4803	5858
SAU203001	4804	5859
SAU203007	4805	5860
SAU203196	4806	5861
SAU203293	4807	5862
SAU203296	4808	5863
SAU203524	4809	5864
SAU300110	4810	5865
SAU300131	4811	5866
SAU300156	4812	5867
SAU300191	4813	5868
SAU300269	4814	5869
SAU300335	4815	5870
SAU300338	4816	5871
SAU300455	4817	5872
SAU300572	4818	5873
SAU300617	4819	5874
SAU300713	4820	5875
SAU300719	4821	5876
SAU300732	4822	5877
SAU300825	4823	5878
SAU300868	4824	5879
SAU300975	4825	5880
SAU300998	4826	5881
SAU301004	4827	5882
SAU301030	4828	5883
SAU301054	4829	5884
SAU301080	4830	5885
SAU301118	4831	5886
SAU301133	4832	5887

PathoSeq Gene Locus	Nucleotide SeqID	Protein SeqID
SAU301148	4833	5888
SAU301223	4834	5889
SAU301230	4835	5890
SAU301268	4836	5891
SAU301275	4837	5892
SAU301357	4838	5893
SAU301363	4839	5894
SAU301433	4840	5895
SAU301465	4841	5896
SAU301472	4842	5897
SAU301592	4843	5898
SAU301620	4844	5899
SAU301758	4845	5900
SAU301773	4846	5901
SAU301829	4847	5902
SAU301869	4848	5903
SAU301898	4849	5904
SAU302060	4850	5905
SAU302513	4851	5906
SAU302626	4852	5907
SAU302685	4853	5908
SAU302698	4854	5909
SAU302699	4855	5910
SAU302805	4856	5911
SAU302901	4857	5912
SAU302931	4858	5913
SAU302950	4859	5914
SAU302956	4860	5915

WHAT IS CLAIMED IS:

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1. A purified or isolated nucleic acid sequence comprising a nucleotide sequence consisting essentially of one of SEQ ID NOs: 8-3795, wherein expression of said nucleic acid inhibits proliferation of a cell.

- 2. A purified or isolated nucleic acid comprising a fragment of one of SEQ ID NOs.: 8-3795, said fragment selected from the group consisting of fragments comprising at least 10, at least 20, at least 25, at least 30, at least 50 and more than 50 consecutive nucleotides of one of SEQ ID NOs: 8-3795.
- 3. A purified or isolated antisense nucleic acid comprising a nucleotide sequence complementary to at least a portion of an intragenic sequence, intergenic sequence, sequences spanning at least a portion of two or more genes, 5' noncoding region, or 3' noncoding region within an operon comprising a proliferation-required gene whose activity or expression is inhibited by an antisense nucleic acid comprising the nucleotide sequence of one of SEO ID NOs.: 8-3795.
- 4. A purified or isolated nucleic acid comprising a nucleotide sequence having at least 70% identity to a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, fragments comprising at least 25 consecutive nucleotides of SEQ ID NOs.: 8-3795, the nucleotide sequences complementary to SEQ ID NOs.: 8-3795 and the sequences complementary to fragments comprising at least 25 consecutive nucleotides of SEQ ID NOs.: 8-3795 as determined using BLASTN version 2.0 with the default parameters.
- 5. A vector comprising a promoter operably linked to a nucleic acid encoding a polypeptide whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence of any one of SEQ ID NOs.: 8-3795.
- 6. A purified or isolated polypeptide comprising a polypeptide whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence of any one of SEQ ID NOs.: 8-3795, or a fragment selected from the group consisting of fragments comprising at least 5, at least 10, at least 20, at least 40, at least 50, at least 60 or more than 60 consecutive amino acids of one of the said polypeptides.
- 7. A purified or isolated polypeptide comprising a polypeptide having at least 25% amino acid identity to a polypeptide whose expression is inhibited by a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, or at least 25% amino acid identity to a fragment comprising at least 10, at least 20, at least 30, at least 40, at least 50, at least 60 or more than 60 consecutive amino acids of a polypeptide whose expression is inhibited by a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 as determined using FASTA version 3.0t78 with the default parameters.
- 8. A method of producing a polypeptide, comprising introducing a vector comprising a promoter operably linked to a nucleic acid comprising a nucleotide sequence encoding a

polypeptide whose expression is inhibited by an antisense nucleic acid comprising one of SEQ ID NOs.: 8-3795 into a cell.

9. A method of inhibiting proliferation of a cell in an individual comprising inhibiting the activity or reducing the amount of a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 or inhibiting the activity or reducing the amount of a nucleic acid encoding said gene product.

10. A method for identifying a compound which influences the activity of a gene product required for proliferation, said gene product comprising a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, said method comprising:

contacting said gene product with a candidate compound; and determining whether said compound influences the activity of said gene product,

- 11. A method for identifying a compound or nucleic acid having the ability to reduce the activity or level of a gene product required for proliferation, said gene product comprising a gene product whose activity or expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, said method comprising:
 - (a) contacting a target gene or RNA encoding said gene product with a candidate compound or nucleic acid; and
 - (b) measuring an activity of said target.

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- 12. A method for identifying a compound which reduces the activity or level of a gene product required for proliferation of a cell, wherein the activity or expression of said gene product is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, said method comprising the steps of:
 - (a) providing a sublethal level of an antisense nucleic acid comprising a nucleotide sequence complementary to a nucleic acid comprising a nucleotide sequence encoding said gene product in a cell to reduce the activity or amount of said gene product in said cell, thereby producing a sensitized cell;
 - (b) contacting said sensitized cell with a compound; and
 - (c) determining the degree to which said compound inhibits proliferation of said sensitized cell relative to a cell which does not contain said antisense nucleic acid.
- 13. A method for inhibiting cellular proliferation comprising introducing an effective amount of a compound with activity against a gene whose activity or expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 or a compound with activity against the product of said gene into a population of cells expressing said gene.

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14. A composition comprising an effective concentration of an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, or a proliferation-inhibiting portion thereof in a pharmaceutically acceptable carrier.

- 15. A method for inhibiting the activity or expression of a gene in an operon required for proliferation wherein the activity or expression of at least one gene in said operon is inhibited by an antisense nucleic acid comprising a sequence selected from the group consisting of SEQ ID NOs.: 8-3795, said method comprising contacting a cell in a cell population with an antisense nucleic acid complementary to at least a portion of said operon.
 - 16. A method for identifying a gene which is required for proliferation of a cell comprising:
 - (a) contacting a cell with an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, wherein said cell is a cell other than the organism from which said nucleic acid was obtained;
 - (b) determining whether said nucleic acid inhibits proliferation of said cell; and
 - (c) identifying the gene in said cell which encodes the mRNA which is complementary to said antisense nucleic acid or a portion thereof.
- 17. A method for identifying a compound having the ability to inhibit proliferation of a cell comprising:
 - (a) identifying a homolog of a gene or gene product whose activity or level is inhibited by a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs. 8-3795 in a test cell, wherein said test cell is not the cell from which said nucleic acid was obtained;
 - (b) identifying an inhibitory nucleic acid sequence which inhibits the activity of said homolog in said test cell;
 - (c) contacting said test cell with a sublethal level of said inhibitory nucleic acid, thus sensitizing said cell:
 - (d) contacting the sensitized cell of step (c) with a compound; and
 - (e) determining the degree to which said compound inhibits proliferation of said sensitized cell relative to a cell which does not contain said inhibitory nucleic acid.
- 18. A method of identifying a compound having the ability to inhibit proliferation comprising:
 - (a) contacting a test cell with a sublethal level of a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs. 8-3795 or a portion thereof which inhibits the proliferation of the cell from which said nucleic acid was obtained, thus sensitizing said test cell;
 - (b) contacting the sensitized test cell of step (a) with a compound; and
 - (c) determining the degree to which said compound inhibits proliferation of said sensitized test cell relative to a cell which does not contain said nucleic acid.

19. A method for identifying a compound having activity against a biological pathway required for proliferation comprising:

- (a) sensitizing a cell by providing a sublethal level of an antisense nucleic acid complementary to a nucleic acid encoding a gene product required for proliferation, wherein the activity or expression of said gene product is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, in said cell to reduce the activity or amount of said gene product;
 - (b) contacting the sensitized cell with a compound; and

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- (c) determining the degree to which said compound inhibits the growth of said sensitized cell relative to a cell which does not contain said antisense nucleic acid.
- 20. A method for identifying a compound having the ability to inhibit cellular proliferation comprising:
 - (a) contacting a cell with an agent which reduces the activity or level of a gene product required for proliferation of said cell, wherein said gene product is a gene product whose activity or expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795;
 - (b) contacting said cell with a compound; and
 - (c) determining whether said compound reduces proliferation of said contacted cell by acting on said gene product.
- 20 21. A method for identifying the biological pathway in which a proliferation-required gene or its gene product lies, wherein said gene or gene product comprises a gene or gene product whose activity or expression is inhibited by an antisense nucleic acid comprising a sequence selected from the group consisting of SEQ ID NOs.: 8-3795, said method comprising:
 - (a) providing a sublethal level of an antisense nucleic acid which inhibits the activity of said proliferation-required gene or gene product in a test cell;
 - (b) contacting said test cell with a compound known to inhibit growth or proliferation of a cell, wherein the biological pathway on which said compound acts is known; and
 - (c) determining the degree to which said proliferation of said test cell is inhibited relative to a cell which was not contacted with said compound.
 - 22. A method for determining the biological pathway on which a test compound acts comprising:
 - (a) providing a sublethal level of an antisense nucleic acid complementary to a proliferation-required nucleic acid in a first cell, wherein the activity or expression of said proliferation-required nucleic acid is inhibited by an antisense nucleic acid comprising a sequence selected from the group consisting of SEQ ID NOs.: 8-3795 and wherein the

biological pathway in which said proliferation-required nucleic acid or a protein encoded by said proliferation-required nucleic acid lies is known,

- (b) contacting said first cell with said test compound; and
- (c) determining the degree to which said test compound inhibits proliferation of said first cell relative to a cell which does not contain said antisense nucleic acid.
- 23. A purified or isolated nucleic acid comprising a sequence selected from the group consisting of SEQ ID NOs.: 8-3795.
- 24. A compound which interacts with a gene or gene product whose activity or expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence of one of SEQ ID NOs.: 8-3795 to inhibit proliferation.
- 25. A compound which interacts with a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence of one of SEQ ID NOs.: 8-3795 to inhibit proliferation.
 - 26. A method for manufacturing an antibiotic comprising the steps of:
- screening one or more candidate compounds to identify a compound that reduces the activity or level of a gene product required for proliferation, said gene product comprising a gene product whose activity or expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795; and

manufacturing the compound so identified.

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- 27. A purified or isolated nucleic acid comprising a nucleic acid having at least 70% nucleotide sequence identity to a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, fragments comprising at least 25 consecutive nucleotides of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, the nucleotide sequences complementary to SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012, and the nucleotide sequences complementary to fragments comprising at least 25 consecutive nucleotides of SEQ ID NOs.: 3796-3800, 3806-4860, 5916-10012 as determined using BLASTN version 2.0 with the default parameters.
- 28. A method of inhibiting proliferation of a cell comprising inhibiting the activity or reducing the amount of a gene product in said cell or inhibiting the activity or reducing the amount of a nucleic acid encoding said gene product in said cell, wherein said gene product is selected from the group consisting of a gene product having having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID

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NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid which hybridizes to a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid which hybridizes to a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs: 8-3795.

29. A method for identifying a compound which influences the activity of a gene product required for proliferation comprising:

contacting a candidate compound with a gene product selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795; and

determining whether said candidate compound influences the activity of said gene product.

- 30. A method for identifying a compound or nucleic acid having the ability to reduce the activity or level of a gene product required for proliferation comprising:
 - (a) providing a target that is a gene or RNA, wherein said target comprises a nucleic acid that encodes a gene product selected from the group consisting of a gene

product having having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleic acid identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEO ID NOs: 8-3795;

- (b) contacting said target with a candidate compound or nucleic acid; and
- (c) measuring an activity of said target.

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31. A method for identifying a compound which reduces the activity or level of a gene product required for proliferation of a cell comprising:

(a) providing a sublethal level of an antisense nucleic acid complementary to a nucleic acid encoding said gene product in a cell to reduce the activity or amount of said gene product in said cell, thereby producing a sensitized cell, wherein said gene product is selected from the group consisting of a gene product having having at least 70% nucleic acid identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs:: 8-3795, a gene product encoded by a nucleic acid comprising a

nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795;

(b) contacting said sensitized cell with a compound; and

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- (c) determining the degree to which said compound inhibits the growth of said sensitized cell relative to a cell which does not contain said antisense nucleic acid.
- 32. A method for inhibiting cellular proliferation comprising introducing a compound with activity against a gene product or a compound with activity against a gene encoding said gene product into a population of cells expressing said gene product, wherein said gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795.
- 33. A preparation comprising an effective concentration of an antisense nucleic acid in a pharmaceutically acceptable carrier wherein said antisense nucleic acid is selected from the group consisting of a nucleic acid comprising a sequence having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 or a proliferation-inhibiting portion thereof, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, and a nucleic acid

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comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795 under moderate conditions.

- 34. A method for inhibiting the activity or expression of a gene in an operon which encodes a gene product required for proliferation comprising contacting a cell in a cell population with an antisense nucleic acid comprising at least a proliferation-inhibiting portion of said operon in an antisense orientation, wherein said gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795. a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEO ID NOs: 8-3795.
 - 35. A method for identifying a gene which is required for proliferation of a cell comprising:
 - (a) contacting a cell with an antisense nucleic acid selected from the group consisting of a nucleic acid at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795 or a proliferation-inhibiting portion thereof, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, wherein said cell is a cell other than the organism from which said nucleic acid was obtained;
 - (b) determining whether said nucleic acid inhibits proliferation of said cell; and
 - (c) identifying the gene in said cell which encodes the mRNA which is complementary to said antisense nucleic acid or a portion thereof.
- 36. A method for identifying a compound having the ability to inhibit proliferation of a cell comprising:

(a) identifying a homolog of a gene or gene product whose activity or level is inhibited by an antisense nucleic acid in a test cell, wherein said test cell is not the microorgaism from which the antisense nucleic acid was obtained, wherein said antisense nucleic acid is selected from the group consisting of a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOs. 8-3795, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795 under moderate conditions;

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- (b) identifying an inhibitory nucleic acid sequence which inhibits the activity of said homolog in said test cell:
- (c) contacting said test cell with a sublethal level of said inhibitory nucleic acid, thus sensitizing said cell;

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- (d) contacting the sensitized cell of step (c) with a compound; and
- (e) determining the degree to which said compound inhibits proliferation of said sensitized cell relative to a cell which does not express said inhibitory nucleic acid.
- 37. A method of identifying a compound having the ability to inhibit proliferation comprising:

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(a) sensitizing a test cell by contacting said test cell with a sublethal level of an antisense nucleic acid, wherein said antisense nucleic acid is selected from the group consisting of a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOs. 8-3795 or a portion thereof which inhibits the proliferation of the cell from which said nucleic acid was obtained, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditionst;

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(b) contacting the sensitized test cell of step (a) with a compound; and

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- (c) determining the degree to which said compound inhibits proliferation of said sensitized test cell relative to a cell which does not contain said antisense nucleic acid.
- 38. A method for identifying a compound having activity against a biological pathway required for proliferation comprising:

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(a) sensitizing a cell by providing a sublethal level of an antisense nucleic acid complementary to a nucleic acid encoding a gene product required for proliferation, wherein said gene product is selected from the group consisting of a gene product having at

least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEO ID NOs: 8-3795;

- (b) contacting the sensitized cell with a compound; and
- (c) determining the extent to which said compound inhibits the growth of said sensitized cell relative to a cell which does not contain said antisense nucleic acid.
 39. A method for identifying a compound having the ability to inhibit cellular proliferation comprising:

(a) contacting a cell with an agent which reduces the activity or level of a gene product required for proliferation of said cell, wherein said gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795

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under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795;

- (b) contacting said cell with a compound; and
- (c) determining the degree to which said compound reduces proliferation of said contacted cell relative to a cell which was not contacted with said agent.
- 40. A method for identifying the biological pathway in which a proliferation-required gene product or a gene encoding a proliferation-required gene product lies comprising:
 - (a) providing a sublethal level of an antisense nucleic acid which inhibits the activity or reduces the level of said gene encoding a proliferation-required gene product or said said proliferation-required gene product in a test cell, wherein said proliferationrequired gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795;
 - (b) contacting said test cell with a compound known to inhibit growth or proliferation of a cell, wherein the biological pathway on which said compound acts is known; and
 - (c) determining the degree to which said compound inhibits proliferation of said test cell relative to a cell which does not contain said antisense nucleic acid.

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41. A method for determining the biological pathway on which a test compound acts comprising:

(a) providing a sublethal level of an antisense nucleic acid complementary to a proliferation-required nucleic acid in a cell, thereby producing a sensitized cell, wherein said antisense nucleic acid is selected from the group consisting of a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795 or a proliferation-inhibiting portion thereof, a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, and a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions and wherein the biological pathway in which said proliferation-required nucleic acid or a protein encoded by said proliferation-required polypeptide lies is known.

(b) contacting said cell with said test compound; and

- (c) determining the degree to which said compound inhibits proliferation of said sensitized cell relative to a cell which does not contain said antisense nucleic acid.
- 42. A compound which inhibits proliferation by interacting with a gene encoding a gene product required for proliferation or with a gene product required for proliferation, wherein said gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.; 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEO ID NOs: 8-3795.
 - 43. A method for manufacturing an antibiotic comprising the steps of:

screening one or more candidate compounds to identify a compound that reduces the activity or level of a gene product required for proliferation wherein said gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795 under moderate conditions, and a gene product whose activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795; and

manufacturing the compound so identified.

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44. A method for inhibiting proliferation of a cell in a subject comprising administering an effective amount of a compound that reduces the activity or level of a gene product required for proliferation of said cell, wherein said gene product is selected from the group consisting of a gene product having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs.: 8-3795, a gene product encoded by a nucleic acid having at least 70% nucleotide sequence identity as determined using BLASTN version 2.0 with the default parameters to a nucleic acid encoding a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NOs:8-3795, a gene product having at least 25% amino acid identity as determined using FASTA version 3.0t78 with the default parameters to a gene product whose expression is inhibited by an antisense nucleic acid comprising a nucleotide sequence selected from the group consisting of SEO ID NOs.: 8-3795, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEQ ID NOs.: 8-3795 under stringent conditions, a gene product encoded by a nucleic acid comprising a nucleotide sequence which hybridizes to a nucleic acid selected from the group consisting of SEO ID NOs.: 8-3795 under moderate conditions, and a gene product whose

activity may be complemented by the gene product whose activity is inhibited by a nucleic acid selected from the group consisting of SEQ ID NOs: 8-3795.

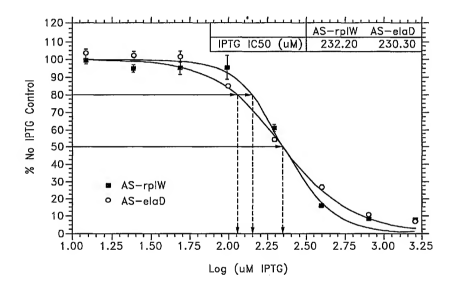


FIG. 1

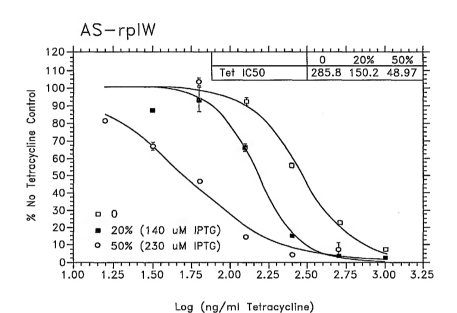


FIG. 2A

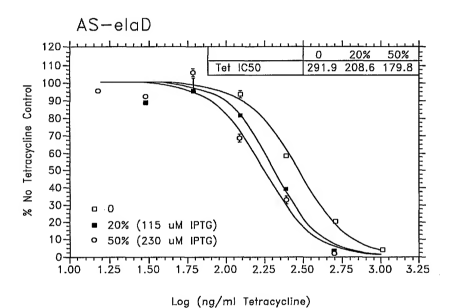


FIG.2B

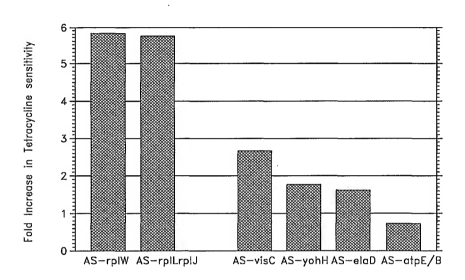
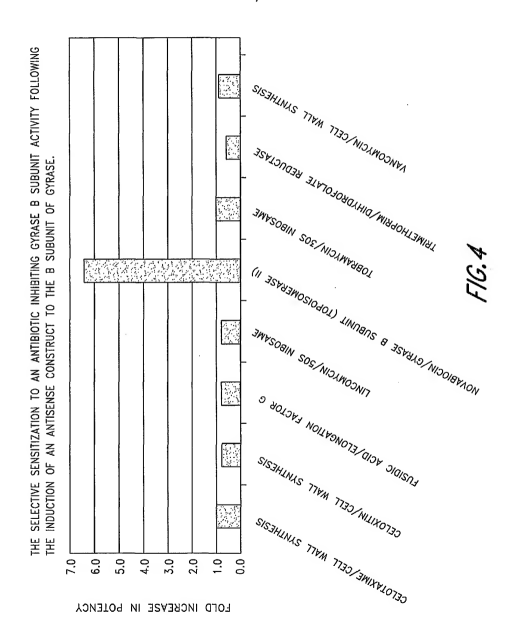


FIG.3



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